



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(54) Title: NOVEL STREPTOCOCCUS ANTIGENS

BVH11-2 SP64	BVH11 SP63	BVH11 JNR.7/87	BVH11-2 JNR.7/87	BVH11 WU2	BVH11-2 WU2	BVH11 A66	BVH11-2 A66	BVH11 P4241	BVH11-2 P4241	BVH11 Rx-1	BVH11-2 Rx-1
181%	183%	188%	182%	180%	180%	180%	180%	180%	180%	183%	181%
S 86%	S 90%	S 91%	S 87%	S 85%	S 85%	S 85%	S 85%	S 85%	S 85%	S 91%	S 85%
187%	187%	198%	195%	196%	195%	196%	195%	195%	196%	187%	194%
S 90%	S 90%	S 98%	S 96%	S 97%	S 96%	S 97%	S 96%	S 97%	S 97%	S 90%	S 95%
196%	188%	188%	187%	188%	187%	187%	188%	187%	197%	189%	BVH11
S 90%	S 91%	S 91%	S 90%	S 91%	S 90%	S 91%	S 91%	S 90%	S 97%	S 91%	SP64
187%	187%	186%	186%	187%	186%	187%	187%	186%	196%	188%	BVH11
S 90%	S 91%	S 91%	S 90%	S 91%	S 90%	S 91%	S 91%	S 90%	S 96%	S 90%	JNR.7/87
196%	197%	196%	197%	196%	197%	196%	197%	197%	187%	194%	BVH11-2
S 97%	S 98%	S 97%	S 98%	S 97%	S 98%	S 97%	S 98%	S 98%	S 90%	S 95%	JNR.7/87
198%	192%	198%	199%	199%	198%	199%	198%	198%	187%	192%	BVH11
S 98%	S 94%	S 98%	S 99%	S 98%	S 99%	S 99%	S 99%	S 98%	S 91%	S 94%	WU2
198%	199%	199%	198%	199%	199%	199%	199%	199%	196%	193%	BVH11-2
S 98%	S 99%	S 99%	S 98%	S 99%	S 99%	S 99%	S 99%	S 99%	S 90%	S 95%	WU2
199%	1100%	199%	199%	199%	199%	199%	199%	197%	192%	192%	BVH11
S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 91%	S 94%	A66	
199%	199%	199%	199%	199%	199%	199%	199%	186%	193%	193%	BVH11-2
S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 90%	S 95%	P4241	
199%	199%	197%	197%	197%	197%	197%	197%	186%	193%	191%	BVH11
S 99%	S 99%	S 91%	S 91%	S 91%	S 91%	S 91%	S 91%	S 90%	S 95%	S 92%	Rx-1

(57) Abstract

Streptococcus proteins and polynucleotides encoding them are disclosed. Said proteins are antigenic and therefore useful vaccine components for the prophylaxis or therapy of streptococcus infection in animals. Also disclosed are recombinant methods of producing the protein antigens as well as diagnostic assays for detecting streptococcus bacterial infection.

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## NOVEL STREPTOCOCCUS ANTIGENS

FIELD OF THE INVENTION

5 The present invention is related to antigens, more particularly protein antigens of streptococcus pneumoniaepathogen which are useful as vaccine components for therapy and/or prophylaxis.

10 BACKGROUND OF THE INVENTION

S. pneumoniae is an important agent of disease in man especially among infants, the elderly and immunocompromised persons. It is a bacterium frequently isolated from 15 patients with invasive diseases such as bacteraemia/septicaemia, pneumonia, meningitis with high morbidity and mortality throughout the world. Even with appropriate antibiotic therapy, pneumococcal infections still result in many deaths. Although the advent of 20 antimicrobial drugs has reduced the overall mortality from pneumococcal disease, the presence of resistant pneumococcal organisms has become a major problem in the world today. Effective pneumococcal vaccines could have a major impact on the morbidity and mortality associated with S. pneumoniae 25 disease. Such vaccines would also potentially be useful to prevent otitis media in infants and young children.

Efforts to develop a pneumococcal vaccine have generally concentrated on generating immune responses to the 30 pneumococcal capsular polysaccharide. More than 80 pneumococcal capsular serotypes have been identified on the basis of antigenic differences. The currently available pneumococcal vaccine, comprising 23 capsular polysaccharides

that most frequently caused disease, has significant shortcomings related primarily to the poor immunogenicity of some capsular polysaccharides, the diversity of the serotypes and the differences in the distribution of 5 serotypes over time, geographic areas and age groups. In particular, the failure of existing vaccines and capsular conjugate vaccines currently in development to protect young children against all serotypes spurs evaluation of other S. pneumoniae components. Although immunogenicity of 10 capsular polysaccharides can be improved, serotype specificity will still represent a major limitation of polysaccharide-based vaccines. The use of a antigenically conserved immunogenic pneumococcal protein antigen, either by itself or in combination with additional components, 15 offers the possibility of a protein-based pneumococcal vaccine.

PCT Publication number WO98/18930 published may 7 1998 20 entitled "Streptococcus Pneumoniae antigens and vaccines" describes certain polypeptides which are claimed to be antigenic. However, no biological activity of these 25 polypeptides is reported.

Therefore there remains an unmet need for Streptococcus 25 antigens that may be used as vaccine components for the prophylaxis and/or therapy of Streptococcus infection.

**SUMMARY OF THE INVENTION**  
30 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55

to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

5 In other aspects, there are provided vectors comprising polynucleotides of the invention operably linked to an expression control region, as well as host cells transfected with said vectors and methods of producing polypeptides comprising culturing said host cells under conditions suitable for expression.

10 In yet another aspect, there are provided novel polypeptides encoded by polynucleotides of the invention.

15 **BRIEF DESCRIPTION OF THE DRAWINGS**

Figure 1 is the DNA sequence of BVH-3 gene; **SEQ ID NO: 1.**

20 Figure 2 is the amino acid sequence of BVH-3 protein; **SEQ ID NO: 2.**

Figure 3 is the DNA sequence of BVH-11 gene; **SEQ ID NO: 3.**

25 Figure 4 is the amino acid sequence of BVH-11 protein; **SEQ ID NO: 4.**

Figure 5 is the DNA sequence of BVH-28 gene; **SEQ ID NO: 5.**

30 Figure 6 is the amino acid sequence of BVH-28 protein; **SEQ ID NO: 6.**

Figure 7 is the DNA sequence of BVH-3A gene which corresponds to the 5' terminal end of BVH-3; **SEQ ID NO: 7.**

Figure 8 is the amino acid sequence of BVH-3A protein; **SEQ ID NO: 8.**

5 Figure 9 is the DNA sequence of BVH-3B gene which corresponds to the 3' terminal end of BVH-3; **SEQ ID NO: 9.**

10 Figure 10 is the amino acid sequence of BVH-3B protein; **SEQ ID NO: 10.**

15 Figure 11 depicts the comparison of the predicted amino acid sequences of the BVH-3 open reading frames from WU2, RX1, JNR.7/87, SP64, P4241 and A66 S. pneumoniae strains by using the program Clustal W from MacVector sequence analysis software (version 6.5). Underneath the alignment, there is a consensus line where \* and . characters indicate identical and similar amino acid residues, respectively.

20 Figure 12 depicts the comparison of the predicted amino acid sequences of the BVH-11 open reading frames from WU2, Rx1, JNR.7/87, SP64, P4241, A66 and SP63 S. pneumoniae strains by using the program Clustal W from MacVector sequence analysis software (version 6.5). Underneath the alignment, there is a consensus line where \* and . characters indicate identical and similar amino acid residues, respectively.

25 Figure 13 depicts the comparison of the predicted amino acid sequences of the BVH-11 proteins from various S. pneumoniae strains. The degrees of identity (I) and similarity (S) were determined by using the program Clustal W from MacVector sequence analysis software (version 6.5).

30 Figure 14 is a DNA sequence containing the complete BVH-3 gene (open reading frame "ORF" at nucleotides 1777 to 4896); **SEQ ID NO: 11.**

Figure 15 is a DNA sequence containing the complete BVH-11 gene (ORF at nucleotides 45 to 2567); **SEQ ID NO: 12.**

5 Figure 16 is a DNA sequence containing the complete BVH-11-2 gene (ORF at nucleotides 114 to 2630); **SEQ ID NO: 13.**

Figure 17 is the amino acid sequence of BVH-11-2 protein; **SEQ ID NO: 14.**

10

Figure 18 is the DNA sequence of SP63 BVH-3 gene; **SEQ ID NO:15.**

15 Figure 19 is the amino acid sequence of SP63 BVH-3 protein; **SEQ ID NO: 16.**

Figure 20 is the amino acid sequence of BVH-3M protein; **SEQ ID NO: 55.**

20 Figure 21 is the amino acid sequence of BVH-3AD protein; **SEQ ID NO: 56.**

Figure 22 is the amino acid sequence of L-BVH-3-AD protein; **SEQ ID NO: 57.**

25

Figure 23 is the amino acid sequence of NEW12 protein; **SEQ ID NO: 58.**

30 Figure 24 is the amino acid sequence of BVH-3C protein; **SEQ ID NO: 59.**

Figure 25 is the amino acid sequence of BVH-11M protein; **SEQ ID NO: 60.**

35 Figure 26 is the amino acid sequence of BVH-11A protein; **SEQ ID NO: 61.**

Figure 27 is the amino acid sequence of BVH-11B (also called New13) protein; **SEQ ID NO: 62.**

5 Figure 28 is the amino acid sequence of BVH-11C protein; **SEQ ID NO: 63.**

Figure 29 is the amino acid sequence of NEW1 protein; **SEQ ID NO: 64.**

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Figure 30 is the amino acid sequence of NEW2 protein; **SEQ ID NO: 65.**

15 Figure 31 is the amino acid sequence of NEW3 protein; **SEQ ID NO: 66.**

Figure 32 is the amino acid sequence of NEW4 protein; **SEQ ID NO: 67.**

20 Figure 33 is the amino acid sequence of NEW5 protein; **SEQ ID NO: 68.**

Figure 34 is the amino acid sequence of NEW6 protein; **SEQ ID NO: 69.**

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Figure 35 is the amino acid sequence of NEW7 protein; **SEQ ID NO: 70.**

30 Figure 36 is the amino acid sequence of NEW8 protein; **SEQ ID NO: 71.**

Figure 37 is the amino acid sequence of NEW9 protein; **SEQ ID NO: 72.**

35 Figure 38 is the amino acid sequence of BVH-11-2M protein; **SEQ ID NO: 73.**

Figure 39 is the amino acid sequence of NEW10 protein; **SEQ ID NO: 74.**

5 Figure 40 is the amino acid sequence of NEW11 protein; **SEQ ID NO: 75.**

Figure 41 is the DNA sequence of NEW12 gene; **SEQ ID NO: 76.**

10 Figure 42 is the amino acid sequence of NEW14 protein; **SEQ ID NO: 77.**

Figure 43 is the amino acid sequence of NEW15 protein; **SEQ ID NO: 78.**

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Figure 44 is the amino acid sequence of NEW16 protein; **SEQ ID NO: 79.**

Figure 45 is the DNA sequence of GBS BVH-71 gene; **SEQ ID NO: 80.**

Figure 46 is the amino acid sequence of GBS BVH-71 protein; **SEQ ID NO: 81.**

25 Figure 47 is the DNA sequence of GAS BVH-71 gene; **SEQ ID NO: 82.**

Figure 48 is the amino acid sequence of GAS BVH-71 protein; **SEQ ID NO: 83.**

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**DETAILED DESCRIPTION OF THE INVENTION**

According to one aspect, the present invention provides an  
35 isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55**

**to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.**

According to one aspect, the present invention provides an  
5 isolated polynucleotide encoding a polypeptide having at least 95% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

10

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 8, 10, 14, 16, 55 to 15 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at 20 least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

25 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79 or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 8, 10, 16, 55, 56, 57, 35 58, 59, 64, 65, 66, 78 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a 5 sequence chosen from SEQ ID NOS: 2, 8, 10, 16, 55, 56, 57, 59, 64, 65, 66, 78 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an 10 isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 4, 14, 58, 60, 61, 62, 63, 67, 68, 69, 70, 71, 72, 73, 74, 75, 77, 79 or fragments, analogs or derivatives thereof.

15 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 4, 14, 60, 61, 62, 63, 67, 20 68, 69, 70, 71, 72, 73, 74, 75, 77, 79 or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an 25 isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 2, 4, 10, 14, 16, 55 to 75, 77 to 79 or fragments, analogs or derivatives thereof.

30 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from SEQ ID NOS: 10, 55 to 75, 77, 78, 79 or fragments, analogs or derivatives thereof.

35 According to one aspect, the present invention provides an

isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence chosen from **SEQ ID NOS: 55 to 75, 77, 78, 79** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from **SEQ ID NOS: 2, 4, 6, 8, 10** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from **SEQ ID NOS: 2, 4, 10, 14, 16** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising a sequence chosen from **SEQ ID NOS: 2, 4, 14, 16** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 2** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 4** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 10** or fragments, analogs or derivatives thereof.

5 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 14** or fragments, analogs or derivatives thereof.

10 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 16** or fragments, analogs or derivatives thereof.

15 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 58** or fragments, analogs or derivatives thereof.

20 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 60** or fragments, analogs or derivatives thereof.

25 According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 62** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 62** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 64** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 67** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 68** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 69** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 72** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 74** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention provides an isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide comprising sequence **SEQ ID NO: 77** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NOS: 2, 4, 6, 8, 10** or fragments, 10 analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 15 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence 20 chosen from **SEQ ID NOS: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence 25 chosen from **SEQ ID NOS: 2, 4, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence 30 chosen from **SEQ ID NOS: 2, 4, 8, 10, 14, 16, 55 to 75, 77 to 79** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence 35 chosen from **SEQ ID NOS: 2, 4, 10, 14, 16, 55 to 75, 77 to 79** or fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOS: 2, 4, 10, 14, 16 or fragments,

5 analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 2 or fragments, analogs or

10 derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 4 or fragments, analogs or

15 derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 10 or fragments, analogs or

20 derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 14 or fragments, analogs or

25 derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence SEQ ID NO: 16 or fragments, analogs or

30 derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from SEQ ID NOS: 10, 55 to 75, 77, 78, 79 or

35 fragments, analogs or derivatives thereof.

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77** or fragments, analogs or derivatives thereof.

10

According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NO: 10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence chosen from **SEQ ID NO: 10, 62, 64, 67, 68, 74, 77** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 58** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 62** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 64** or fragments, analogs or derivatives thereof.

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According to one aspect, the present invention relates to

polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 67** or fragments, analogs or derivatives thereof.

- 5 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 68** or fragments, analogs or derivatives thereof.
- 10 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 74** or fragments, analogs or derivatives thereof.
- 15 According to one aspect, the present invention relates to polypeptides characterized by the amino acid sequence comprising sequence **SEQ ID NO: 77** or fragments, analogs or derivatives thereof.
- 20 In a further embodiment, the present invention also relates to chimeric polypeptides which comprise one or more polypeptides or fragments, analogs or derivatives thereof as described in the present application.
- 25 In a further embodiment, the present invention also relates to chimeric polypeptides which comprise one or more polypeptides or fragments, analogs or derivatives thereof as defined in the figures of the present application.
- 30 In a further embodiment, the present application also relates to chimeric polypeptides which comprise two or more polypeptides chosen from **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof ;provided that the polypeptides or
- 35 fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from **SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77** or 5 fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

10 In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from **SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 74, 77** or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

15 In a further embodiment, the chimeric polypeptide will comprise two or more polypeptides chosen from **SEQ ID NOS :10, 62, 64, 67, 68, 74, 77** or fragments, analogs or derivatives thereof; provided that the polypeptides or 20 fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.

In a further embodiment, the chimeric polypeptide will comprise between 2 and 5 polypeptides.

25 In a further embodiment, the chimeric polypeptide will comprise between 2 and 4 polypeptides.

30 In a further embodiment, the chimeric polypeptide will comprise between 2 and 3 polypeptides.

In a further embodiment, the chimeric polypeptide will comprise 2 polypeptides.

In a further embodiment, there is provided a chimeric polypeptide of formula (I):

**A-(B)<sub>m</sub>-(C)<sub>n</sub>-D (I)**

5 Wherein;

**m** is 0 or 1,

**n** is 0 or 1,

10 **A** is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;

**B** is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;

15 **C** is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; and

**D** is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

20

In a further embodiment,

**A** is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof;

25 **B** is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77, or fragments, analogs or derivatives thereof;

**C** is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives

30 thereof; and

**D** is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

35 In a further embodiment,

A is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof;

B is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 74, 77, or fragments, analogs or derivatives thereof;

5 C is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof; and

D is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

10 In one embodiment, chimeric polypeptides of the present invention comprise those wherein the following embodiments are present, either independently or in combination.

In a further embodiment, A is SEQ ID NOS :10, 58, 62, 64, 15 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :10 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :58 or fragments, 20 analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :62 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

25 In a further embodiment, A is SEQ ID NO :67 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :68 or fragments, analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :74 or fragments, 30 analogs or derivatives thereof.

In a further embodiment, A is SEQ ID NO :77 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NOS :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :10 or fragments, 5 analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

10 In a further embodiment, **B** is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :67 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :68 or fragments, 15 analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO :74 or fragments, analogs or derivatives thereof.

In a further embodiment, **B** is SEQ ID NO : 77 or fragments, analogs or derivatives thereof.

20 In a further embodiment, **C** is SEQ ID NOS :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO :10 or fragments, 25 analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO : 62 or fragments, analogs or derivatives thereof.

30 In a further embodiment, **C** is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO : 67 or fragments, analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO : 68 or fragments,

analogs or derivatives thereof.

In a further embodiment, **C** is SEQ ID NO : 74 or fragments, analogs or derivatives thereof.

5 In a further embodiment, **C** is SEQ ID NO : 77 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is SEQ ID NO :10, 58, 62, 64, 67, 68, 74, 77 or fragments, analogs or derivatives thereof.

10 In a further embodiment, **D** is SEQ ID NO :10 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is SEQ ID NO :58 or fragments, analogs or derivatives thereof.

15 In a further embodiment, **D** is SEQ ID NO :62 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is SEQ ID NO :64 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is SEQ ID NO :67 or fragments, analogs or derivatives thereof.

20 In a further embodiment, **D** is SEQ ID NO :68 or fragments, analogs or derivatives thereof.

In a further embodiment, **D** is SEQ ID NO :74 or fragments, analogs or derivatives thereof.

25 In a further embodiment, **D** is SEQ ID NO :77 or fragments, analogs or derivatives thereof.

In a further embodiment, **m** is 0.

In a further embodiment, **n** is 0.

30

In a further embodiment, **m** and **n** are 0.

In a further embodiment, **m** and **n** are 0, **A** is SEQ ID NO:64 or fragments, analogs or derivatives thereof, **B** is SEQ ID

**NO:62** or fragments, analogs or derivatives thereof.

In a further embodiment, **m** and **n** are 0, **A** is **SEQ ID NO:62** or fragments, analogs or derivatives thereof, **B** is **SEQ ID NO:64** or fragments, analogs or derivatives thereof.

5

In accordance with the present invention, all nucleotides encoding polypeptides and chimeric polypeptides are within the scope of the present invention.

10 In a further embodiment, the polypeptides or chimeric polypeptides in accordance with the present invention are antigenic.

15 In a further embodiment, the polypeptides or chimeric polypeptides in accordance with the present invention can elicit an immune response in an individual.

20 In a further embodiment, the present invention also relates to polypeptides which are able to raise antibodies having binding specificity to the polypeptides or chimeric polypeptides of the present invention as defined above.

25 An antibody that "has binding specificity" is an antibody that recognizes and binds the selected polypeptide but which does not substantially recognize and bind other molecules in a sample, e.g., a biological sample, which naturally includes the selected peptide. Specific binding can be measured using an ELISA assay in which the selected polypeptide is used as an antigen.

30

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. All publications, patent applications, patents, and other references mentioned herein are incorporated by reference in their entirety. In

case of conflict, the present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

5

As used herein, "fragments", "derivatives" or "analogs" of the polypeptides of the invention include those polypeptides in which one or more of the amino acid residues are substituted with a conserved or non-conserved 10 amino acid residue (preferably conserved) and which may be natural or unnatural. In one embodiment, derivatives and analogs of polypeptides of the invention will have about 70% identity with those sequences illustrated in the figures or fragments thereof. That is, 70% of the residues 15 are the same. In a further embodiment, polypeptides will have greater than 75% homology. In a further embodiment, polypeptides will have greater than 80% homology. In a further embodiment, polypeptides will have greater than 85% homology. In a further embodiment, polypeptides will have 20 greater than 90% homology. In a further embodiment, polypeptides will have greater than 95% homology. In a further embodiment, polypeptides will have greater than 99% homology. In a further embodiment, derivatives and analogs 25 of polypeptides of the invention will have fewer than about 20 amino acid residue substitutions, modifications or deletions and more preferably less than 10. Preferred substitutions are those known in the art as conserved i.e. the substituted residues share physical or chemical properties such as hydrophobicity, size, charge or 30 functional groups.

In accordance with the present invention, polypeptides of the invention include both polypeptides and chimeric polypeptides.

35

Also included are polypeptides which have fused thereto

other compounds which alter the polypeptides biological or pharmacological properties i.e. polyethylene glycol (PEG) to increase half-life; leader or secretory amino acid sequences for ease of purification; prepro- and pro- sequences; and (poly)saccharides.

Furthermore, in those situations where amino acid regions are found to be polymorphic, it may be desirable to vary one or more particular amino acids to more effectively 10 mimic the different epitopes of the different streptococcus strains.

Moreover, the polypeptides of the present invention can be modified by terminal -NH<sub>2</sub> acylation (eg. by acetylation, or 15 thioglycolic acid amidation, terminal carboxy amidation, e.g. with ammonia or methylamine) to provide stability, increased hydrophobicity for linking or binding to a support or other molecule.

20 Also contemplated are hetero and homo polypeptide multimers of the polypeptide fragments, analogues and derivatives. These polymeric forms include, for example, one or more polypeptides that have been cross-linked with cross-linkers such as avidin/biotin, gluteraldehyde or dimethyl- 25 superimide. Such polymeric forms also include polypeptides containing two or more tandem or inverted contiguous sequences, produced from multicistronic mRNAs generated by recombinant DNA technology. Preferably, a fragment, analog or derivative of a 30 polypeptide of the invention will comprise at least one antigenic region i.e. at least one epitope.

In order to achieve the formation of antigenic polymers (i.e. synthetic multimers), polypeptides may be utilized 35 having bishaloacetyl groups, nitroarylhalides, or the like, where the reagents being specific for thio groups.

Therefore, the link between two mercapto groups of the different peptides may be a single bond or may be composed of a linking group of at least two, typically at least four, and not more than 16, but usually not more than about 5 14 carbon atoms.

In a particular embodiment, polypeptide fragments, analogs and derivatives of the invention do not contain a methionine (Met) starting residue. Preferably, 10 polypeptides will not incorporate a leader or secretory sequence (signal sequence). The signal portion of a polypeptide of the invention may be determined according to established molecular biological techniques. In general, the polypeptide of interest may be isolated from a 15 streptococcus culture and subsequently sequenced to determine the initial residue of the mature protein and therefore the sequence of the mature polypeptide.

According to another aspect, there are provided vaccine 20 compositions comprising one or more streptococcus polypeptides of the invention in admixture with a pharmaceutically acceptable carrier diluent or adjuvant. Suitable adjuvants include oils i.e. Freund's complete or incomplete adjuvant; salts i.e.  $AlK(SO_4)_2$ ,  $AlNa(SO_4)_2$ , 25  $AlNH_4(SO_4)_2$ , silica, kaolin, carbon polynucleotides i.e. poly IC and poly AU. Preferred adjuvants include QuilA and Alhydrogel. Vaccines of the invention may be administered parenterally by injection, rapid infusion, nasopharyngeal absorption, dermoabsorption, or bucal or oral. 30 Pharmaceutically acceptable carriers also include tetanus toxoid.

Vaccine compositions of the invention are used for the treatment or prophylaxis of streptococcus infection and/or 35 diseases and symptoms mediated by streptococcus infection as described in P.R. Murray (Ed, in chief), E.J. Baron, M.A.

Pfaller, F.C. Tenover and R.H. Yolken. Manual of Clinical Microbiology, ASM Press, Washington, D.C. sixth edition, 1995, 1482p which are herein incorporated by reference. In one embodiment, vaccine compositions of the present

5 invention are used for the treatment or prophylaxis of meningitis, otitis media, bacteremia or pneumonia. In one embodiment, vaccine compositions of the invention are used for the treatment or prophylaxis of *streptococcus* infection and/or diseases and symptoms mediated by *streptococcus*

10 infection, in particular *S.pneumoniae*, group A *streptococcus* (*pyogenes*), group B *streptococcus* (GBS or *agalactiae*), *dysgalactiae*, *uberis*, *nocardia* as well as *Staphylococcus aureus*. In a further embodiment, the *streptococcus* infection is *S.pneumoniae*.

15

In a particular embodiment, vaccines are administered to those individuals at risk of *streptococcus* infection such as infants, elderly and immunocompromised individuals.

20 As used in the present application, the term "individuals" include mammals. In a further embodiment, the mammal is human.

Vaccine compositions are preferably in unit dosage form of

25 about 0.001 to 100  $\mu$ g/kg (antigen/body weight) and more preferably 0.01 to 10  $\mu$ g/kg and most preferably 0.1 to 1  $\mu$ g/kg 1 to 3 times with an interval of about 1 to 6 week intervals between immunizations.

30 According to another aspect, there are provided polynucleotides encoding polypeptides characterized by the amino acid sequence chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

35

In one embodiment, polynucleotides are those illustrated in **SEQ ID Nos: 1, 3, 5, 7, 9, 11, 12, 13, 15, 76, 80, 82** which may include the open reading frames (ORF), encoding polypeptides of the invention. It will be appreciated that

5 the polynucleotide sequences illustrated in the figures may be altered with degenerate codons yet still encode the polypeptides of the invention. Accordingly the present invention further provides polynucleotides which hybridize to the polynucleotide sequences herein above described (or

10 the complement sequences thereof) having 50% identity between sequences. In one embodiment, at least 70% identity between sequences. In one embodiment, at least 75% identity between sequences. In one embodiment, at least 80% identity between sequences. In one embodiment, at least 85% identity

15 between sequences. In one embodiment, at least 90% identity between sequences. In a further embodiment, polynucleotides are hybridizable under stringent conditions i.e. having at least 95% identity. In a further embodiment, more than 97% identity.

20

In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 3, 7, 9, 11, 12, 13, 15, 76, 80, 82** encoding polypeptides of the invention.

25 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 3, 9, 11, 12, 13, 15, 76, 80, 82** which may include the open reading frames (ORF), encoding polypeptides of the invention.

30 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 3, 9, 11, 12, 13, 15, 76** which may include the open reading frames (ORF), encoding polypeptides of the invention.

35 In a further embodiment, polynucleotides are those

illustrated in **SEQ ID NOS : 1, 3, 7, 9, 11, 12, 13, 15, 76** which may include the open reading frames (ORF), encoding polypeptides of the invention.

- 5 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 7, 9, 11, 15, 76** which may include the open reading frames (ORF), encoding polypeptides of the invention.
- 10 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 9, 11, 15, 76** which may include the open reading frames (ORF), encoding polypeptides of the invention.
- 15 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 1, 7, 9, 11** which may include the open reading frames (ORF), encoding polypeptides of the invention.
- 20 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO : 1**, encoding polypeptides of the invention.  
In a further embodiment, polynucleotides are those
- 25 illustrated in **SEQ ID NO :7**, encoding polypeptides of the invention.  
In a further embodiment, polynucleotides are those
- 30 illustrated in **SEQ ID NO :9**, encoding polypeptides of the invention.  
In a further embodiment, polynucleotides are those

In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :15**, encoding polypeptides of the invention.

5 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NOS : 3, 12, 13, 76**, encoding polypeptides of the invention.

10 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :3**, encoding polypeptides of the invention.

15 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :12**, encoding polypeptides of the invention.

20 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :13**, encoding polypeptides of the invention.

25 In a further embodiment, polynucleotides are those illustrated in **SEQ ID NO :76**, encoding polypeptides of the invention.

30 As will be readily appreciated by one skilled in the art, polynucleotides include both DNA and RNA.

35 The present invention also includes polynucleotides complementary to the polynucleotides described in the present application.

In a further aspect, polynucleotides encoding polypeptides of the invention, or fragments, analogs or derivatives thereof, may be used in a DNA immunization method. That is, they can be incorporated into a vector which is

replicable and expressible upon injection thereby producing the antigenic polypeptide in vivo. For example polynucleotides may be incorporated into a plasmid vector under the control of the CMV promoter which is functional 5 in eukaryotic cells. Preferably the vector is injected intramuscularly.

According to another aspect, there is provided a process for producing polypeptides of the invention by recombinant 10 techniques by expressing a polynucleotide encoding said polypeptide in a host cell and recovering the expressed polypeptide product. Alternatively, the polypeptides can be produced according to established synthetic chemical techniques i.e. solution phase or solid phase synthesis of 15 oligopeptides which are ligated to produce the full polypeptide (block ligation).

General methods for obtention and evaluation of polynucleotides and polypeptides are described in the 20 following references: Sambrook et al, Molecular Cloning: A Laboratory Manual, 2nd ed, Cold Spring Harbor, N.Y., 1989; Current Protocols in Molecular Biology, Edited by Ausubel F.M. et al., John Wiley and Sons, Inc. New York; PCR Cloning Protocols, from Molecular Cloning to Genetic 25 Engineering, Edited by White B.A., Humana Press, Totowa, New Jersey, 1997, 490 pages; Protein Purification, Principles and Practices, Scopes R.K., Springer-Verlag, New York, 3rd Edition, 1993, 380 pages; Current Protocols in Immunology, Edited by Coligan J.E. et al., John Wiley & 30 Sons Inc., New York which are herein incorporated by reference.

For recombinant production, host cells are transfected with vectors which encode the polypeptide, and then cultured in 35 a nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the genes.

Suitable vectors are those that are viable and replicable in the chosen host and include chromosomal, non-chromosomal and synthetic DNA sequences e.g. bacterial plasmids, phage DNA, baculovirus, yeast plasmids, vectors derived from 5 combinations of plasmids and phage DNA. The polypeptide sequence may be incorporated in the vector at the appropriate site using restriction enzymes such that it is operably linked to an expression control region comprising a promoter, ribosome binding site (consensus region or 10 Shine-Dalgarno sequence), and optionally an operator (control element). One can select individual components of the expression control region that are appropriate for a given host and vector according to established molecular biology principles (Sambrook et al, Molecular Cloning: A 15 Laboratory Manual, 2nd ed, Cold Spring Harbor, N.Y., 1989; Current Protocols in Molecular Biology, Edited by Ausubel F.M. et al., John Wiley and Sons, Inc. New York incorporated herein by reference). Suitable promoters include but are not limited to LTR or SV40 promoter, *E.coli* 20 lac, tac or trp promoters and the phage lambda  $P_l$  promoter. Vectors will preferably incorporate an origin of replication as well as selection markers i.e. ampicillin resistance gene. Suitable bacterial vectors include pET, pQE70, pQE60, pQE-9, pbs, pD10 phagescript, psiX174, 25 pbluescript SK, pbsks, pNH8A, pNH16a, pNH18A, pNH46A, ptrc99a, pKK223-3, pKK233-3, pDR540, pRIT5 and eukaryotic vectors pBlueBacIII, pWLNEO, pSV2CAT, pOG44, pXT1, pSG, pSVK3, pBPV, pMSG and pSVL. Host cells may be bacterial i.e. *E.coli*, *Bacillus subtilis*, *Streptomyces*; fungal i.e. 30 *Aspergillus niger*, *Aspergillus nidulans*; yeast i.e. *Saccharomyces* or eukaryotic i.e. CHO, COS.

Upon expression of the polypeptide in culture, cells are typically harvested by centrifugation then disrupted by 35 physical or chemical means (if the expressed polypeptide is not secreted into the media) and the resulting crude

extract retained to isolate the polypeptide of interest. Purification of the polypeptide from culture media or lysate may be achieved by established techniques depending on the properties of the polypeptide i.e. using ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, hydroxylapatite chromatography and lectin chromatography. Final purification may be achieved using HPLC.

10

The polypeptide may be expressed with or without a leader or secretion sequence. In the former case the leader may be removed using post-translational processing (see US 4,431,739; US 4,425,437; and US 4,338,397 incorporated herein by reference) or be chemically removed subsequent to purifying the expressed polypeptide.

According to a further aspect, the streptococcus polypeptides of the invention may be used in a diagnostic test for streptococcus infection, in particular S. pneumoniae infection. Several diagnostic methods are possible, for example detecting streptococcus organism in a biological sample, the following procedure may be followed:

- a) obtaining a biological sample from a patient;
- 25 b) incubating an antibody or fragment thereof reactive with a streptococcus polypeptide of the invention with the biological sample to form a mixture; and
- c) detecting specifically bound antibody or bound fragment in the mixture which indicates the presence

30 of streptococcus.

Alternatively, a method for the detection of antibody specific to a streptococcus antigen in a biological sample containing or suspected of containing said antibody may be performed as follows:

- a) obtaining a biological sample from a patient;

- b) incubating one or more streptococcus polypeptides of the invention or fragments thereof with the biological sample to form a mixture; and
- c) detecting specifically bound antigen or bound fragment in the mixture which indicates the presence of antibody specific to streptococcus.

5 One of skill in the art will recognize that this diagnostic test may take several forms, including an immunological

10 10 test such as an enzyme-linked immunosorbent assay (ELISA), a radioimmunoassay or a latex agglutination assay, essentially to determine whether antibodies specific for the protein are present in an organism.

15 15 The DNA sequences encoding polypeptides of the invention may also be used to design DNA probes for use in detecting the presence of streptococcus in a biological sample suspected of containing such bacteria. The detection method of this invention comprises:

20 20 a) obtaining the biological sample from a patient;

b) incubating one or more DNA probes having a DNA sequence encoding a polypeptide of the invention or fragments thereof with the biological sample to form a mixture; and

25 25 c) detecting specifically bound DNA probe in the mixture which indicates the presence of streptococcus bacteria.

30 30 The DNA probes of this invention may also be used for detecting circulating streptococcus i.e.

S.pneumoniaenucleic acids in a sample, for example using a polymerase chain reaction, as a method of diagnosing streptococcus infections. The probe may be synthesized using conventional techniques and may be immobilized on a

35 35 solid phase, or may be labelled with a detectable label. A preferred DNA probe for this application is an oligomer

having a sequence complementary to at least about 6 contiguous nucleotides of the streptococcus pneumoniae polypeptides of the invention.

5 Another diagnostic method for the detection of streptococcus in a patient comprises:

- a) labelling an antibody reactive with a polypeptide of the invention or fragment thereof with a detectable label;
- 10 b) administering the labelled antibody or labelled fragment to the patient; and
- c) detecting specifically bound labelled antibody or labelled fragment in the patient which indicates the presence of streptococcus.

15

A further aspect of the invention is the use of the streptococcus polypeptides of the invention as immunogens for the production of specific antibodies for the diagnosis and in particular the treatment of streptococcus infection.

20 Suitable antibodies may be determined using appropriate screening methods, for example by measuring the ability of a particular antibody to passively protect against streptococcus infection in a test model. One example of an animal model is the mouse model described in the examples

25 herein. The antibody may be a whole antibody or an antigen-binding fragment thereof and may belong to any immunoglobulin class. The antibody or fragment may be of animal origin, specifically of mammalian origin and more specifically of murine, rat or human origin. It may be a

30 natural antibody or a fragment thereof, or if desired, a recombinant antibody or antibody fragment. The term recombinant antibody or antibody fragment means antibody or antibody fragment which was produced using molecular biology techniques. The antibody or antibody fragments may

35 be polyclonal, or preferably monoclonal. It may be specific for a number of epitopes associated with the

streptococcus pneumoniae polypeptides but is preferably specific for one.

Without limiting its scope, the present invention also 5 relates to new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The present invention also relates to truncated polypeptides comprising fragments of the new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The present invention also relates to chimeric 10 polypeptides comprising fragments of the new antigens designated BVH-3, BVH-11, BVH-11-2, BVH-28 and BVH-71. The following is a reference table summarizing the relation between the antigens of the present invention:

Family	Nucleotide SEQ ID NO	Polypeptide SEQ ID NO
<b>BVH-3</b>		
BVH-3	1, 11	2
BVH-3A	7	8
BVH-3B	9	10
BVH-3 SP63	15	16
BVH-3M		55
BVH-3AD		56
L-BVH-3AD		57
New12	76	58
BVH-3C		59
New1		64
New2		65
New3		66
New15		78
<b>BVH-11</b>		
BVH-11	3, 12	4
BVH-11-2	13	14
BVH-11M		60
BVH-11A		61
BVH-11B also referred to as NEW13		62
BVH-11C		63
New4		67
New5		68

Family	Nucleotide SEQ ID NO	Polypeptide SEQ ID NO
New6		69
New7		70
New8		71
New9		72
BVH-11-2M		73
New10		74
New11		75
New12	76	58
New14		77
New16		79
BVH-28		
BVH-28	5	6
BVH-71		
GBS	80	81
GAS	82	83

## EXAMPLE 1

5 This example illustrates the cloning of S. pneumoniae genes.

The coding region of S. pneumoniae gene BVH-3 (SEQ ID NO: 1) and the coding region of S. pneumoniae gene BVH-28 (SEQ ID NO: 5) were amplified by PCR (DNA Thermal Cycler GeneAmp 10 PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of serogroup 6 S. pneumoniae strain SP64 using the oligos that contained base extensions for the addition of restriction sites BglII (AGATCT) and XbaI (TCTAGA). PCR products were purified from agarose gel using a QIAquick gel extraction kit from QIAGen (Chatsworth, CA), digested BglII-XbaI (Pharmacia Canada Inc, Baie d'Urfé, Canada), extracted with phenol : chloroform and precipitated with ethanol. The Superlinker vector pSL301 (Invitrogen, San Diego, CA) was digested with BglII and XbaI and purified from agarose gel 15 using a QIAquick gel extraction kit from QIAGen (Chatsworth, CA). The BglII-XbaI genomic DNA fragments were ligated to 20

the BglIII-XbaI pSL301 vector. The ligated products were transformed into E. coli strain DH5a [f80 lacZ DM15 endA1 recA1 hsdR17 (<sup>r</sup>K<sup>m</sup>K<sup>r</sup>) supE44 thi-11<sup>r</sup> gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the 5 method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). Recombinant pSL301 plasmids (rpSL301) containing either BVH-3 or BVH-28 gene were purified using a QIAgen kit (Chatsworth, CA) and DNA inserts were confirmed by nucleotide sequence analysis (Taq Dye 10 Deoxy Terminator Cycle Sequencing kit, ABI, Foster City, CA). Recombinant rpSL301 (rpSL301) were digested with the restriction enzymes BglII (AGATCT) and XhoI (CTCGAG). DNA fragments BglII-XhoI were purified using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). pET-32c(+) 15 expression vector (Novagen, Madison, WI) containing the thioredoxin-His·Tag sequence was digested with BamHI (GGATCC) and XhoI and gel extracted using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The BglII-XhoI DNA fragments were ligated to the BamHI-XhoI pET-32c(+) 20 vector to create the coding sequence for thioredoxin-His·Tag-BVH-3 or thioredoxin-His·Tag-BVH-28 fusion protein. The ligated products were transformed into E. coli strain DH5a [f80 lacZ DM15 endA1 recA1 hsdR17 (<sup>r</sup>K<sup>m</sup>K<sup>r</sup>) supE44 thi-11<sup>r</sup> gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, 25 MD) according to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). Recombinant pET-32c(+) plasmids were purified using a QIAgen kit (Chatsworth, CA) and the nucleotide sequences at the fusion sites of thioredoxin-His·Tag and DNA insert were verified by 30 DNA sequencing (Taq Dye Deoxy Terminator Cycle Sequencing kit, ABI, Foster City, CA).

## EXAMPLE 2

5 This example illustrates the cloning of S. pneumoniae protein genes in CMV plasmid pCMV-GH.

The DNA coding region of a S. pneumoniae protein was inserted in phase downstream of a human growth hormone (hGH) gene which was under the transcriptional control of the 10 cytomegalavirus (CMV) promotor in the plasmid vector pCMV-GH (Tang et al., Nature, 1992, 356 :152). The CMV promotor is non functional plasmid in E. coli cells but active upon administration of the plasmid in eukaryotic cells. The vector also incorporated the ampicillin resistance gene.

15 The coding region of BVH-3 gene (**SEQ ID NO: 1**) and BVH-28 gene (**SEQ ID NO: 5**) were obtained from rpSL301 (see example 1) using restriction enzymes BglII (AGATCT) and XbaI (TCTAGA). The digested products were purified from agarose 20 gel using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The pCMV-GH vector (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas) containing the human growth hormone to create fusion proteins was digested with 25 BglII and XbaI and purified from agarose gel using the QIAquick gel extraction kit from QIAgen (Chatsworth, CA). The BglII-XbaI DNA fragments were ligated to the BglII-XbaI pCMV-GH vector to create the hGH-BVH-3 or hGH-BVH-28 fusion protein under the control of the CMV promoter. The ligated 30 products were transformed into E. coli strain DH5a[f80 lacZ DM15 endA1 recA1 hsdR17 (<sup>r</sup>K<sup>m</sup>K<sup>r</sup>) supE44 thi-11<sup>r</sup> gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according

to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). The recombinant pCMV plasmids were purified using a QIAgen kit (QIAgen, Chatsworth, CA).

5

The coding region of BVH-11 gene (**SEQ ID NO: 3**) was amplified by PCR (DNA Thermal Cycler GeneAmp PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of serogroup 6 S. pneumoniae strain SP64 using the oligos that contained 10 base extensions for the addition of restriction sites BglII (AGATCT) and HindIII (AAGCTT). The PCR product was purified from agarose gel using a QIAquick gel extraction kit from QIAgen (Chatsworth, CA), digested with restriction enzymes (Pharmacia Canada Inc, Baie d'Urfe, Canada), extracted with 15 phenol : chloroform and precipitated with ethanol. The pCMV-GH vector (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas) was digested with BglII and HindIII and purified from agarose gel using the QIAquick gel extraction kit from 20 QIAgen (Chatsworth, CA). The BglII-HindIII DNA fragment was ligated to the BglII-HindIII pCMV-GH vector to create the hGH-BVH-11 fusion protein under the control of the CMV promoter. The ligated products were transformed into E. coli strain DH5a[f80 lacZ DM15 endA1 recA1 hsdR17 ( $\lambda$ K<sup>M</sup>K<sup>S</sup>) supE44 25 thi-11<sup>r</sup> gyrA96 relA1 D(lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, D. DNA Cloning, 1985, D.M. Glover (ed), pp. 109-135). The recombinant pCMV plasmid was purified using a QIAgen kit (Chatsworth, CA) and the nucleotide sequence of 30 the DNA insert was verified by DNA sequencing.

## EXAMPLE 3

This example illustrates the use of DNA to elicit an immune response to S. pneumoniae antigens.

5

A group of 8 female BALB/c mice (Charles River, St-Constant, Québec, Canada) were immunized by intramuscular injection of 50 µl three times at two- or three-week intervals with 100 µg of recombinant pCMV-GH encoding the BVH-3, BVH-11 or the 10 BVH-28 gene in presence of 50 µg of granulocyte-macrophage colony-stimulating factor (GM-CSF)- expressing plasmid pCMV-GH-GM-CSF (Laboratory of Dr. Stephen A. Johnston, Department of Biochemistry, The University of Texas, Dallas, Texas). As control, a group of mice were injected with 100 µg of 15 pCMV-GH in presence of 50 µg of pCMV-GH-GM-CSF. Blood samples were collected from the orbital prior to each immunization and seven days following the third injection and serum antibody responses were determined by ELISA using thioredoxin-His·Tag-S. pneumoniae fusion protein as coating 20 antigen. DNA immunization with recombinant plasmid pCMV-GH encoding the BVH-3, BVH-11 or the BVH-28 S. pneumoniae protein induced antibody reactive against the respective recombinant protein. The reciprocal antibody titers, defined as the highest serum dilution at which the absorbance values 25 were 0.1 above the background values, were above  $4 \times 10^3$ .

## EXAMPLE 4

30 This example illustrates the production and purification of recombinant S. pneumoniae proteins.

The recombinant pET plasmids containing the BVH-3, BVH-11 or the BVH-28 gene corresponding to the **SEQ ID NO: 1**, **SEQ ID NO: 3** or the **SEQ ID NO: 5** respectively were transformed by

5 electroporation (Gene Pulser II apparatus, BIO-RAD Labs, Mississauga, Canada) into E. coli strain AD494 (DE3) (Dara-leu7697 DlacX74 DphoA PvuII phoR DmalF3 F' [lac<sup>r</sup>(lacI<sup>q</sup>) pro] trxB::Kan) (Novagen, Madison, WI). In this strain of E. coli, the T7 promotor controlling expression of the fusion

10 protein is specifically recognized by the T7 RNA polymerase (present on the 1DE3 prophage) whose gene is under the control of the lac promotor which is inducible by isopropyl-β-d-thio-galactopyranoside (IPTG). The transformant AD494(DE3)/rpET was grown at 37°C with agitation at 250 rpm

15 in LB broth (peptone 10g/L, yeast extract 5g/L, NaCl 10g/L) containing 100μg of ampicillin (Sigma-Aldrich Canada Ltd., Oakville, Canada) per ml until the A<sub>600</sub> reached a value of 0.6. In order to induce the production of the thioredoxin-His·Tag-BVH-3, thioredoxin-His·Tag-BVH-11 or thioredoxin-

20 His·Tag-BVH-28 fusion protein, the cells were incubated for 2 additional hours in the presence of IPTG at a final concentration of 1 mM. Induced cells from a 100 ml culture were pelleted by centrifugation and frozen at -70°C.

25 The purification of the fusion proteins from the soluble cytoplasmic fraction of IPTG-induced AD494(DE3)/rpET was done by affinity chromatography based on the properties of the His·Tag sequence (6 consecutive histidine residues) to bind to divalent cations (Ni<sup>2+</sup>) immobilized on the His·Bind

30 metal chelation resin. Briefly, the pelleted cells obtained from a 100mL culture induced with IPTG were resuspended in

phosphate-buffered (PBS):500mM NaCl pH7.1, sonicated and spun at 20,000 X g for 20 min to remove debris. The supernatant was filtered (0.22 $\mu$ m pore size membrane) and deposited on a HiTrap® 1mL chelating pre-packed ready-to-use 5 column (Pharmacia Biotech, Baie d'Urfé, Canada). The thioredoxin-His·Tag-S. pneumoniae fusion protein was eluted with 1M imidazole-500mM NaCl-PBS pH7.1. The removal of the salt and imidazole from the sample was done by dialysis against PBS at 4°C. The quantities of fusion protein 10 obtained from the soluble fraction of E. coli was estimated by MicroBCA (Pierce, Rockford, Illinois).

#### EXAMPLE 5

15

This example illustrates the protection of mice against fatal pneumococcal infection by immunization.

Groups of 8 female BALB/c mice (Charles River) were 20 immunized subcutaneously three times at three-week intervals with either 25  $\mu$ g of affinity purified thioredoxin-His·Tag-BVH-3 fusion protein in presence of 15  $\mu$ g of QuilA adjuvant (Cedarlane Laboratories Ltd, Hornby, Canada) or, as control, with QuilA adjuvant alone in PBS. Blood samples were 25 collected from the orbital sinus on day 1, 22 and 43 prior to each immunization and seven days (day 50) following the third injection. One week later the mice were challenged with approximately  $10^6$  CFU of the type 3 S. pneumoniae strain WU2. Samples of the S. pneumoniae challenge inoculum 30 were plated on chocolate agar plates to determine the CFU and to verify the challenge dose. Deaths were recorded for

a period of 14 days and on day 14 post-challenge, the surviving mice were sacrificed and blood samples tested for the presence of S. pneumoniae organisms. The survival data are shown in table 1.

5

Prechallenge sera were analyzed for the presence of antibodies reactive with S. pneumoniae by standard immunoassays. Elisa and immunoblot analyses indicated that immunization with recombinant S. pneumoniae protein produced 10 in E. coli elicited antibodies reactive with both, recombinant and native pneumococcal protein.

Table 1. Protection mediated by recombinant BVH-3 protein

Immunogen	No. of mice alive : no. of mice dead 14 days post-challenge	Median day of death
BVH-3	8 : 0	>14
none	0 : 8	1

15

All mice immunized with BVH-3 recombinant protein survived to infection while none of the control mice given adjuvant alone survived. There was a significant difference in survival between the two groups of mice ( $P<0.0001$ , log rank 20 test for nonparametric analysis of survival curves;  $P=0.0002$ , Fisher's exact test). All hemocultures from surviving mice were negative at day 14 post-challenge.

25

EXAMPLE 6

This example describes the cloning of BVH-3 and BVH-11 genes from a variety of S. pneumoniae strains and the molecular conservation of these genes.

5 Molecular analysis of chromosomal DNA from various S. pneumoniae isolates with DNA probes spanning different regions of BVH-3 or BVH-11 revealed the presence of one BVH-3 gene copy and two BVH-11 gene copies. The two BVH-11 gene copies are not identical and the genes were  
10 arbitrarily designated BVH-11 (SEQ ID NO:12; ORF at nucleotides 45 to 2567) and BVH-11-2 (SEQ ID NO:13; ORF at nucleotides 114 to 2630).

15 The first amino acids of the BVH-3 and BVH-11 coding regions have the characteristics of leader sequences also known as signal peptides. The consensus signal peptidase cleavage site L-X-X-C of lipoprotein modification/processing sites was present in the sequences. Mature BVH-3, BVH-11 and BVH-11-2 proteins from S. pneumoniae SP64 have 1019, 821 and 819 amino acids, respectively. The regions of S. pneumoniae genes coding for mature BVH-3, termed BVH-3M, (nucleotides 1837 - 4896; SEQ. ID. NO: 11), BVH-11M (nucleotides 102-2567; SEQ. ID. NO: 12) and BVH-11-2M (nucleotides 171-2630; SEQ. ID. NO: 13), were amplified by PCR(DNA Thermal Cycler GeneAmp PCR system 2400 Perkin Elmer, San Jose, CA) from genomic DNA of 25 6 or 7 S. pneumoniae strains. Serogroup 6 S. pneumoniae SP64 and serogroup 9 SP63 clinical isolates were provided by the laboratoire de la santé publique du Québec, Sainte-  
30 Anne-de-Bellevue; serotype 4 strain JNR.7/87 was provided by Andrew Camilli, Tufts University School of Medicine, Boston; Rx1 strain, a nonencapsulated derivative of the type 2 strain D39 and the type 3 strains A66 and WU2 were provided by David E. Briles from University of Alabama, Birmingham and the type 3 clinical isolate P4241 was provided by the centre de recherche en infectiologie du

centre hospitalier de l'université Laval, Sainte-Foy. The sets of oligonucleotide primers OCRR479-OCRR480; HAMJ160-OCRR488 and HAMJ160-HAMJ186, that contained base extensions for the addition of restriction sites were used for the 5 amplification of BVH-3, BVH-11 and BVH-11-2 gene, respectively, with the exception of BVH-11 gene from SP64 strain which was amplified using the set of primers consisting of HAMJ487 and OCRR488. Primer sequences are listed below (Table 2). PCR products were purified from 10 agarose gel using a QIAquick gel extraction kit from QIAGen (Chatsworth, CA) and digested BglII-XbaI or BglII-HindIII (Pharmacia Canada Inc, Baie d'Urfé, Canada). Digestions were cleaned using a QIAquick PCR purification kit from QIAGen (Chatsworth, CA). The PCR products were ligated to 15 the BglII-XbaI or BglII-HindIII pSL301 vector. The ligated products were transformed into E. coli strain DH5 $\alpha$  [ $\phi$ 80 lacZ  $\Delta$ M15 endA1 recA1 hsdR17 ( $^r$ K $^m$ K $^s$ ) supE44 thi-1 $\lambda$  gyrA96 relA1  $\Delta$ (lacZYA-argF)U169] (Gibco BRL, Gaithersburg, MD) according to the method of Simanis (Hanahan, D. DNA 20 Cloning, 1985, D.M. Glover (ed), pp. 109-135). Recombinant pSL301 plasmids (rpSL301) containing BVH-3, BVH-11 or BVH11-2 were purified using a QIAGen kit (Chatsworth, CA) and DNA inserts were sequenced (Taq Dye Deoxy Terminator Cycle Sequencing kit, ABI, Foster City, CA). The figures 11 25 and 12 depict the consensus sequence established from the BVH-3, and BVH-11 deduced amino acid sequences, respectively. Comparison of BVH-3 protein sequences revealed 99 to 100% identity of sequences for all strains with the exception that BVH-3 from serogroup 9 SP63 strain 30 (SEQ. ID. NO: 15 and SEQ. ID. NO: 16) misses a stretch of 177 amino acids corresponding to residues 244 to 420 on BVH-3' protein sequence of S. pneumoniae SP64. Analysis of sequences of additional serogroup 9 strains revealed BVH-3 molecule having the same deletion in 3 out of 4 strains

thus suggesting that the 3 strains are members of a S. pneumoniae serogroup 9 clone.

Comparison of 13 BVH-11 nucleotide sequences obtained from 5 7 S. pneumoniae strains, revealed that the nucleotide sequences are very similar. Computer analysis (MacVector, Clustal W 1.4) using multiple alignment of the predicted BVH-11 protein sequences revealed that these sequences were 10 75% identical and 82 % homologous on a length of 834 amino acids. Pairwise alignment revealed 80 to 100% identity (Figure 13). The sequences showed great similarity in overall organization. Variability in the primary sequence of these proteins is almost restricted to the last 125 15 amino acids in the C-terminal portion of the proteins. This region constitutes a domain. Close examination of this domain revealed two groups of sequences. The first 9 sequences from the figure 13 belong to one group while the last 4 sequences belong to another group. A 39% identity value is obtained when the domain sequences of the 13 20 proteins are compared (MacVector, Clustal W 1.4). The identity value increased to more than 92% when sequences belonging to a same group are compared.

25 EXAMPLE 7

This example illustrates the homology of portions of BVH-3 and BVH-11 genes.

30 Molecular analysis with DNA probes derived from BVH-3 and BVH-11 genes indicated that BVH-3 and BVH-11 were related. In dot blot hybridization studies, DNA probe consisting of either, BVH-3 or BVH-11, gene sequence hybridized to both, BVH-3 and BVH-11 genes thus indicating that BVH-3 and BVH-35 11 genes shared homologous sequences. Comparison of sequences revealed that the ORFs and the proteins were 43

and 33% identical, respectively. Closer examination revealed that the region corresponding to amino acids 1 to 225 in BVH-3 and 1 to 228 in BVH-11 were 73 and 75% identical at the DNA and protein level, respectively. In 5 contrast, the 3' regions corresponding to amino acids 226 to 1039 from BVH-3 and amino acids 229-840 from BVH-11 were only 34 and 22% identical at the DNA and protein level, respectively. Thus the 5' termini of BVH-3 and BVH-11 genes appear to contain highly conserved sequences while 10 the remaining parts of the genes are highly divergent. These results suggest that BVH-3 and BVH-11 might share similar functions mediated by sequences present in the conserved region whereas BVH-3- and BVH-11-specific functions might be mediated by sequences in the divergent 15 region.

#### EXAMPLE 8

20 This example describes the cloning of truncated BVH-3, BVH-11 and BVH-11-2 genes by polymerase chain reaction (PCR) and the expression of truncated BVH-3 and BVH-11 molecules.

25 Gene fragments were amplified by PCR using pairs of oligonucleotide engineered to amplify fragments spanning the BVH-3 (**SEQ ID NO: 1** and **SEQ ID NO: 11**), BVH-11 (**SEQ ID NO: 3** and **SEQ ID NO: 12**) or BVH-11-2 (**SEQ ID NO: 13**) gene from S. pneumoniae strain SP64. Each of the primers had a restriction endonuclease site at the 5' end, thereby 30 allowing directional in-frame cloning of the amplified product into the digested plasmid vector (Tables 2 and 3). PCR-amplified products were digested with restriction endonucleases and ligated to either linearized plasmid pSL301 (see example 1), pCMV-GH (see example 2) or pET 35 (Novagen, Madison, WI) expression vector digested likewise or digested with enzymes that produce compatible cohesive

ends. Recombinant pSL301 and recombinant pCMV-GH plasmids were digested with restriction enzymes for the in-frame cloning in pET expression vector. Clones were first stabilized in E. coli DH5 $\alpha$  before introduction into E. coli BL21( $\lambda$ DE3) or AD494 ( $\lambda$ DE3) for expression of truncated BVH-3 or BVH-11 molecules. Each of the resultant plasmid constructs was confirmed by nucleotide sequence analysis. The recombinant proteins were expressed as N-terminal fusions with the thioredoxin and His-tag or as C-terminal fusions with an His-tag. The expressed recombinant proteins were purified from supernatant fractions obtained from centrifugation of sonicated IPTG-induced E. coli cultures using a His-Bind metal chelation resin (QIAgen, Chatsworth, CA). The gene products generated are listed in the table 3. The gene products corresponding to the N-terminal region including the signal sequence are designated as Lippedated-proteins or lipoproteins (L-proteins). The gene products corresponding to the N-terminal region lacking the signal sequence are identified as protein without signal sequence (w/o ss).

Table 2. List of PCR oligonucleotide primers

Primer	SEQ. ID.	Sequence 5' - 3'	Nucleotide position	Restriction sites
OCRR 479	17	cagtagatctgtgcctatgcactaac	SEQ ID 1:61-78	BglII
OCRR 480	18	gatctctagactactgtctattccgtacgtatg	SEQ ID 11:4909-4887	XbaI
OCRR 497	19	atcactcgaggattacctggataatccgt	SEQ ID 1:1525-1506	XhoI
OCRR 498	20	ctgctaaggcttatgaaagatttagat	SEQ ID 1:1534-1548	HindIII

OCRR 499	21	gatactcgagctgctattccctac	SEQ ID 11 :4906-4893	XhoI
HAMJ 172	22	gaatctcgagtaagctgctgtaattc	SEQ ID 1 : 675-661	XhoI
HAMJ 247	23	gacgctcgagcgctatgaaatcagataaattc	SEQ ID 1 :3117-3096	XhoI
HAMJ 248	24	gacgctcgagggcattaccctggataatcctgttcatg	SEQ ID 1 :1527-1501	XhoI
HAMJ 249	25	cagtagatcttcatcatttattgaaaagagg	SEQ ID 11 : 1749-1771	BglII
HAMJ 278	26	ttatttctccatatggacttgacagaagagcaaattaag	SEQ ID 1 :1414-1437	NdeI
HAMJ 279	27	cgcacaagctcgctatgaaatcagataaattc	SEQ ID 1 :3117-3096	HindIII
HAMJ 280	28	cgcacaagctttccacaatataaagtgcattgatt	SEQ ID 1 :2400-2377	HindIII
HAMJ 281	29	ttatttctccatatggaaagtacctatcttggaaaaagaa	SEQ ID 1 :2398-2421	NdeI
HAMJ 300	30	ttatttctccatatgggcctatgcactaaaccagc	SEQ ID 1 :62-82	NdeI
HAMJ 313	31	ataagaatgcggccgttccacaatataaagtgcattgatt	SEQ ID 1 :2400-2377	NotI
OCRR 487	32	cagtagatctgtgcattgaacttaggttgc	SEQ ID 3 :58-79	BglII
OCRR 488	33	gatcaagctgtgcacctttacttactctc	SEQ ID 12 :2577-2556	HindIII
HAMJ 171	34	ctgagatatccgttatcgtaaacc	SEQ ID 3 :1060-1075	EcoRV
HAMJ 251	35	ctgcaagctttaaagggaaataatcgc	SEQ ID 3 :1059-1045	HindIII
HAMJ 264	36	cagtagatctgcagaaggcattctatctg	SEQ ID 3 :682-700	BglII
HAMJ 282	37	tcgccaagcttcgttatcgtaaaccattggg	SEQ ID 3 :1060-1081	HindIII
HAMJ 283	38	ataagaatgcggccgcctactctccttaataaagccaatagtt	SEQ ID 3 :2520-2492	NdeI
HAMJ 284	39	catgccatggacattgatagtctcttgcacagc	SEQ ID 3 :856-880	NcoI
HAMJ 285	40	cgcacaagcttacttcctttaataaagccaatag	SEQ ID 3 :2520-2494	HindIII
HAMJ 286	41	cgcacaagcttaacatggcgtacgttacc	SEQ ID 3 :2139-2119	HindIII
HAMJ 287	42	cataccatggccattatgaggcacctaag	SEQ ID 3 :2014-2034	NcoI
HAMJ 288	43	cgcacaagcttaagtaatcttcagcctctcag	SEQ ID 3 :2376-2353	HindIII

HAMJ 289	44	gataccatggctagcgaccatgtcaaagaa	SEQ ID 3:2125-2146	NcoI
HAMJ 290	45	cgc caagcttatcatccactaactgactttatcac	SEQ ID 3:1533-1508	HindIII
HAMJ 291	46	cataccatggatattctgcctcttagctccg	SEQ ID 3:1531-1554	NcoI
HAMJ 301	47	catgccatggtgcttatgaacttaggtttgc	SEQ ID 3:59-79	NcoI
HAMJ 302	48	cgc caagcttagcggttacccaaaaccattatc	SEQ ID 3:2128-2107	HindIII
HAMJ 160	49	gtatttagatctgttcctatgaacttggcgtcacca	SEQ ID 13: 172-196	BglII
HAMJ 186	50	cgcctcttagactactgtataggagccgg	SEQ ID 13: 2460-2443	XbaI
HAMJ 292	51	catgccatggaaaacatttcaagccctttacgtg	SEQ ID 11: 754-778	NcoI
HAMJ 293	52	cgc aagcttctgtataggagccggttgactttc	SEQ ID 11: 2457-2434	HindIII
HAMJ 294	53	catgccatgggttcgtaaaaataaggcagaccaag	SEQ ID 11: 2038-2062	NcoI
HAMJ 297	54	catgccatggaaaggctattggaatggaaag	SEQ ID 11: 622-642	NcoI

Table 3. Lists of truncated BVH-3 and BVH-11 gene products generated from S. pneumoniae SP64

PCR-primer sets	Protein designation	Identification (encoded amino acids)	SEQ. ID.NO.	Cloning vector
OCRR479-OCRR480	BVH-3M	BVH-3 w/o ss (21-1039)	55	pSL301
OCRR479-OCRR497	BVH-3AD	BVH-3 N' end w/o ss (21-509)	56	pSL301
HAMJ248-HAMJ249	L-BVH-3AD	BVH-3 N' end (1-509)	57	pET-21(+)
OCRR498-OCRR499	BVH-3B	BVH-3 C' end (512-1039)	10	pSL301
OCRR479-HAMJ172	BVH-3C	BVH-3 N' end w/o ss (21-225)	59	pET-32 c(+)
OCRR487-OCRR488	BVH-11M	BVH-11 w/o ss (20-840)	60	pCMV-GH
HAMJ251-OCRR487	BVH-11A	BVH-11 N' end w/o ss (20-353)	61	pET-32 c(+)
HAMJ171-OCRR488	BVH-11B	BVH-11 C' end (354-840)	62	pET-32 a(+)
HAMJ264-OCRR488	BVH-11C	BVH-11 C' end (228-840)	63	pET-32 a(+)
HAMJ278-HAMJ279	NEW1	BVH-3 C' end (472-1039)	64	pET-21b(+)
HAMJ278-HAMJ280	NEW2	BVH-3 C' end (472-800)	65	pET-21b(+)
HAMJ281-HAMJ279	NEW3	BVH-3 C' end (800-1039)	66	pET-21b(+)
HAMJ284-HAMJ285	NEW4	BVH-11 C' end (286-840)	67	pET-21d(+)
HAMJ284-HAMJ286	NEW5	BVH-11 internal (286-713)	68	pET-21d(+)
HAMJ287-HAMJ288	NEW6	BVH-11 internal (672-792)	69	pET-21d(+)
HAMJ285-HAMJ289	NEW7	BVH-11 internal (709-840)	70	pET-21d(+)
HAMJ284-HAMJ290	NEW8	BVH-11 internal (286-511)	71	pET-21d(+)

HAMJ286-HAMJ291	NEW9	BVH-11 internal (511-713)	72	pET-21d(+)
HAMJ160-HAMJ186	BVH-11-2M	BVH-11-2 w/o ss (20-838)	73	pSL301
HAMJ292-HAMJ293	NEW10	BVH-11-2 C' end (271-838)	74	pET-21d(+)
HAMJ293-HAMJ294	NEW11	BVH-11-2 C' end (699-838)	75	pET-21d(+)
HAMJ282-HAMJ283	BVH-11B	BVH-11 C' end (354-840)	62	pET-21b(+)
HAMJ286-HAMJ297	NEW14	BVH-11-2 internal (227-699)	77	pET-21d(+)
HAMJ300-HAMJ313	NEW15	BVH-3 N' end w/o ss (21-800)	78	pET-21b(+)
HAMJ301-HAMJ302	NEW16	BVH-11 N' end w/o ss (20-709)	79	pET-21d(+)

## EXAMPLE 9

This example describes the isolation of monoclonal antibodies (Mabs) and the use of Mabs to characterize BVH-3, BVH-11 and BVH-11-2 protein epitopes.

Female BALB/c mice (Charles River) were immunized subcutaneously with BVH-3, BVH-11 or BVH-11-2 gene products from S. pneumoniae strain SP64 in presence of 15 µg of QuilA adjuvant (Cedarlane Laboratories Ltd, Hornby, Canada). One set of mice (fusion experiment 1) were immunized on day 1 and 14 with 25 µg of affinity purified thioredoxin-His•Tag-BVH-3M fusion protein. A second group of mice (fusion experiment 2) were immunized three times at three-week intervals with 25 µg of affinity purified thioredoxin-His•Tag-BVH-11M. A third group of mice (fusion experiment 3) were immunized on day 1 and day 15 with 25 µg of affinity purified thioredoxin-His•Tag-BVH-11-2M fusion protein. A fourth group of mice (fusion experiment 4) were immunized on day 1 with 25 µg of affinity purified thioredoxin-His•BVH-11B fusion protein and boosted by intravenous injection on day 16 and on day 37 with recombinant BVH-11B in PBS. Three to four days before fusion, mice were injected intravenously with 25 µg of the respective antigen suspended in PBS alone. Hybridomas were produced by fusion of spleen cells with nonsecreting SP2/0 myeloma cells as previously described by J. Hamel et al. [J. Med. Microbiol., 23, pp163-170 (1987)]. Culture supernatants of hybridomas were initially screened by enzyme-linked-immunoassay according to the procedure described by Hamel et al. (Supra) using plates coated with preparations of purified recombinant proteins or suspensions of heat-killed S. pneumoniae cells. Positive hybridomas selected on the basis of ELISA reactivity with a

variety of antigens were then cloned by limiting dilutions, expanded and frozen.

Hybridomas were tested by ELISA or Western immunoblotting  
 5 against BVH-3 and BVH-11 gene products in order to characterize the epitopes recognized by the Mabs. BVH-3 and BVH-11 shared common epitopes with 6 Mabs (H3-1-F9, H3-1-D4, H3-1-H12, H11-1-E7, H11-1-H10 and H11-1.1-G11) showing reactivities with both proteins (Table 4). BVH-11  
 10 and BVH-11-2 molecules from S. pneumoniae SP64 shared common epitopes not present on BVH-3 with Mabs (3A1, 13C11, 10H10, 1D8, 10G9, 10A2, 3E8, 10D7, 2H7 and 6H7) reactive with both, BVH-11 and BVH-11-2, recombinant proteins (Table 5).

15

Table 4. Reactivity of BVH-3-immunoreactive Mabs with a panel of BVH-3 and BVH-11 gene products

MAbs	a. Immunoreactivity with						
	BVH-3M 21-1039	BVH-3A 21-509	BVH-3B 512-1039	BVH-3C 21-225	NEW2 472-800	NEW3 800-1039	BVH-11M 20-840
H3-1-F9	+	+	-	+	-	-	+
H3-1-D4	+	+	-	+	-	-	+
H3-1-H12	+	+	-	+	-	-	+
H3-2-G2	+	+	-	-	-	-	-
H3-3-A1	+	+	-	-	-	-	-
H3-4-D3	+	-	+	-	-	+	-
H11-1-E7	+	+	-	+	-	-	+
H11-1- H10	+	+	-	+	-	-	+
H11- 1.1-G11	+	+	-	+	+	-	+

Table 5. Reactivity of Mabs raised against BVH-11-2 protein from S. pneumoniae strain SP64 with a panel of BVH-11 gene products

Mabs*	b. Immunoreactivity with							
	c. BVH-11 products				d. BVH-11-2 products			
	BVH-11M 20-840	NEW8 286-511	NEW9 511-713	BVH-11B 354-840	BVH-11-2 20-838	NEW10 271-838	NEW11 699-838	NEW14 227-699
3A1	+	+	-	+	+	+	-	+
13C1	+	+	+	+	+	+	-	+
10H10	+	+	+	+	+	+	-	+
1D8	+	+	-	+	+	+	-	+
10G9	+	-	-	+	+	+	-	+
10A2	+	-	-	+	+	+	-	+
3E8	+	-	-	+	+	+	-	+
10D7	+	-	-	+	+	+	-	+
2H7	+	-	-	-	+	-	-	-
6H7	+	-	-	-	+	-	-	-
3A4	-	-	-	-	+	+	+	-
14H6	-	-	-	-	+	+	+	-
7G2	-	-	-	-	+	+	-	+
13H10	-	-	-	-	+	-	-	+
7E8	-	-	-	-	+	-	-	-
7H6	-	-	-	-	+	-	-	-

\* Mabs listed in this table were not reactive with recombinant BVH-3 molecule

5

The results obtained from the immunoreactivity studies of the Mabs (Table 4 and Table 5) are in agreement with the protein sequences derived from the respective gene sequences. Indeed the Mabs cross-reactive with BVH-3 and BVH-11 molecules recognized BVH-3C protein corresponding to the conserved region, and BVH-11 and BVH-11-2 specific Mabs were reactive with epitopes located on variable parts of these molecules. BVH-3 and BVH-11, and BVH-11 and BVH-11-2 can be distinguished by their reactivity with Mabs.

15

#### EXAMPLE 10

This example illustrates the simultaneous expression of BVH-3 and BVH-11 gene products by S. pneumoniae.

A standard Western blot technique was used to investigate  
5 whether BVH-3 and BVH-11 genes were expressed in S.  
pneumoniae. S. pneumoniae strain SP64 and SP63 were grown  
overnight at 37°C in 5% CO<sub>2</sub> on chocolate agar plates,  
bacteria were suspended in PBS and heat-killed at 56°C for  
20 min. For the preparation of antigens, suspensions of S.  
10 pneumoniae were treated with sample buffer containing SDS  
and 2-mercaptoethanol for 5 min at 100°C. Pneumococcal  
protein antigens were resolved by SDS-PAGE electrophoresis  
according to the method of Laemmli [Nature, 227, pp. 680-  
685 (1970)]. After SDS-PAGE, the proteins were transferred  
15 electrophoretically from the gel to nitrocellulose paper by  
the method of Towbin [Proc. Natl. Acad. Sci. USA, 76, pp.  
4350-4354 (1979)] and probed with mouse antiserum or  
monoclonal antibodies. The detection of antigens reactive  
with the antibodies was performed by indirect enzyme-  
20 immunoassay using conjugated-anti-mouse immunoglobulins and  
a colour substrate. When antiserum raised to recombinant  
BVH-3 was tested against S. pneumoniae SP64 antigens, two  
reactive bands having apparent molecular masses of 127 kDa  
and 99 kDa were detected. Bands having the same apparent  
25 molecular masses were also detected when Mabs H3-1-F9, H3-  
1-D4, H3-1-H12, H11-1-E7, H11-1-H10 and H11-1.1-G11 were  
used individually as immunological probes. In contrast,  
Mabs specific for the BVH-3 molecule detected the 127 kDa  
band only and Mabs specific for BVH-11 detected the 99 kDa  
30 band only thus confirming the identity of the 127 and 99  
kDa bands as BVH-3 and BVH-11, respectively. These studies  
provide evidence that BVH-3 and BVH-11 proteins are  
simultaneously present on S. pneumoniae. Moreover, the  
results are consistent with our previous observations that  
35 BVH-3 and BVH-11 possess epitopes that are common to both  
proteins and epitopes that are exclusive to either protein.

In S. pneumoniae SP64, mature BVH-3, BVH-11 and BVH-11-2 are proteins of 1019, 821 and 819 amino acids with predicted molecular mass of 112.5 kDa, 92.4 kDa, and 91.7 kDa, respectively. Although there is a discrepancy between the molecular mass predicted from the sequence and the molecular mass calculated on SDS-PAGE, BVH-3 can be distinguished from BVH-11 by its higher molecular mass. Moreover, BVH-3 molecules from S. pneumoniae strain SP63 have an apparent molecular mass of 112 kDa in SDS-PAGE compared to 127 kDa for BVH-3 of SP64 strain. This data is consistent with the deletion of a stretch of 177 amino acid residues in BVH-3 of S. pneumoniae strain SP63.

15

## EXAMPLE 11

This example describes the protection conferred in experimental infection of mice vaccinated with recombinant BVH-3 or BVH-11 gene products.

Groups of 7 or 8 female BALB/c mice (Charles River) were immunized subcutaneously three times at three-week intervals with either affinity purified thioredoxin-25 His•Tag-BVH-3M fusion protein, affinity purified thioredoxin-His•Tag-BVH-11M fusion protein or, as control, with QuilA adjuvant alone in PBS. Twelve to 14 days following the third immunization, the mice were challenged intravenously with S. pneumoniae WU2 strain or intranasally 30 with P4241 strain. Samples of the S. pneumoniae challenge inoculum were plated on chocolate agar plates to determine the CFU and to verify the challenge dose. The challenge dose was approximately  $10^6$  CFU. Deaths were recorded for a period of 14 days and on day 14 post-challenge, the 35 surviving mice were sacrificed and blood samples tested for

the presence of S. pneumoniae organisms. The survival data are shown in Tables 6 and 7.

5

Table 6. Protection mediated by recombinant BVH-3M and BVH-11M proteins in experimental infection with virulent S. pneumoniae WU2

Experiment	Immunogen	Alive : dead <sup>a</sup>	Median days alive
1	BVH-3M	8 : 0	>14
	none	0 : 8	1
2	BVH-11M	8 : 0	>14
	none	0 : 8	1

10 <sup>a</sup> The number of mice alive : the number of mice dead on day 14 post-challenge.

15 Table 7. Protection mediated by recombinant BVH-3M and BVH-11M proteins in experimental pneumonia with virulent S. pneumoniae P4241

Experiment	Immunogen	Alive : dead <sup>a</sup>	Median day alive
1	BVH-3M	6 : 1	>14
	none	1 : 7	4.5
2	BVH-3M	8 : 0	>14
	BVH-11M	8 : 0	>14
	none	0 : 8	4

16 <sup>a</sup> The number of mice alive : the number of mice dead on day 14 post-challenge.

17 All mice immunized with recombinant BVH-3M or BVH-11M  
 20 protein survived to infection with WU2 while none of the control mice given adjuvant alone survived. All except one mice immunized with recombinant BVH-3M or BVH-11M protein survived to infection with P4241 while only one control mice given adjuvant alone survived. All hemocultures from

surviving mice were negative at day 14 post-challenge. These results clearly indicate that both, BVH-3M and BVH-11M, elicit protective anti-pneumococcal immune responses in mice. The fact that these proteins are highly conserved 5 among S. pneumoniae isolates emphasize the potential of BVH-3 and BVH-11 as universal vaccine candidates. Indeed, the BVH-3 and BVH-11 proteins from serogroup 6 S. pneumoniae strain SP64 elicited protection against pneumococcal infections with strains of different capsular 10 serotypes.

Ideally, a vaccine that could protect against pneumococcal disease, could protect against meningitis, otitis media, bacteremia and pneumonia. BVH-3 and BVH-11 were protective 15 against lethal systemic- and pneumonia-infection models thus suggesting that, in humans, BVH-3- and BVH11-protein-based vaccines could reduce the incidence of a wide spectrum of disease caused by virtually all S. pneumoniae independently of the capsular serotype.

20 Data from Tables 6 and 7 clearly demonstrate that BVH-3 and BVH-11 were, both, protection-eliciting molecules of S. pneumoniae. It was not known, however, whether protection can be mediated by specific sequences that were not shared 25 on BVH-3 and BVH-11 molecules. Groups of female BALB/c mice (Charles River) were immunized subcutaneously three times at three-week intervals with either affinity purified thioredoxin-His•Tag- BVH-3AD, -BVH-3B or -BVH-3C fusion protein in presence of 15 µg of QuilA adjuvant (Cedarlane 30 Laboratories Ltd, Hornby, Canada). Control mice were immunized with QuilA adjuvant alone in PBS or affinity purified thioredoxin-His•Tag or thioredoxin-His•Tag-fusion protein (His-Thio) in presence of QuilA.

35 To determine the protective ability of a set of truncated proteins, termed NEW4, NEW5, NEW6, NEW7, NEW8, NEW9, NEW10,

NEW11, NEW14 and BVH-11B, groups of female BALB/c mice (Charles River) were immunized subcutaneously two times at three-week intervals with 25 µg of either affinity purified His•Tag-fusion protein in presence of 15 µg of Quila adjuvant. Ten to 14 days following the last immunization, the mice were challenged with virulent S. pneumoniae. Our results indicate that, BVH-3B, a truncated BVH-3 molecule consisting of amino acids 512-1039, elicited protection against the mouse-virulent strains WU2 and P4241.

Similarly, BVH-11B, NEW4 and NEW5 molecules, three truncated BVH-11 molecules consisting of amino acids 354-840, amino acids 286-840 and amino acids 286-713, respectively, elicited protection against experiment intravenous challenge with WU2 and intranasal challenge with P4241. Moreover, vaccination with NEW10 and NEW14, consisting of amino acids 272-838 and amino acids 227-699 from BVH-11-2 molecule also resulted in protection against death with the pneumococcal strains. These results indicate that the region comprising 428 amino acids extending from amino acids 286-713 and amino acids 272-699 on S. pneumoniae SP64 BVH-11 and BVH-11-2 protein sequences, respectively, contains protective epitopes. This region is highly conserved with a global 91% identity and 94% homology among thirteen BVH-11 protein sequences.

25

Table 8. Evaluation of protection elicited by vaccination of mice with BVH-3 and BVH-11 gene products

Experiment	Immunogen	Challenge with WU2		Challenge with P4241	
		Alive : dead <sup>a</sup>	Median day alive	Alive : dead	Median day alive
1 <sup>b</sup>	None	0 : 8	1.5	1 : 7	4.5
	NEW4	8 : 0	>14	8 : 0	>14
	NEW5	8 : 0	>14	8 : 0	>14
	NEW7	0 : 8	2	0 : 8	5
	BVH-11M	8 : 0	>14	8 : 0	>14
2 <sup>b</sup>	None	0 : 8	1	0 : 8	4
	NEW5	8 : 0	>14	8 : 0	>14
	NEW8	0 : 8	1.5	0 : 8	5.5
	NEW9	3 : 5	3.5	2 : 6	7
	BVH-11M	8 : 0	>14	8 : 0	>14
3 <sup>b</sup>	None	0 : 8	1	0 : 8	4
	NEW6	0 : 8	1	4 : 4	10.5 <sup>c</sup>
	NEW10	8 : 0	>14	8 : 0	>14
	NEW11	0 : 8	1.5	1 : 7	6
	BVH-11M	8 : 0	>14	8 : 0	>14
4 <sup>b</sup>	None	0 : 8	2	0 : 8	4
	BVH-11B	7 : 1	>14	8 : 0	>14
	NEW14	8 : 0	>14	8 : 0	>14
5	His-Thio	0 : 8	2		
	BVH-3AD	1 : 7	2.5		
	BVH-3B	5 : 3	>14		
6	His-Thio	0 : 8	1		
	BVH-3C	0 : 8	1		

<sup>a</sup> The number of mice alive : the number of mice dead on day 14 post-challenge.

<sup>b</sup> The WU2 challenge dose was 10<sup>5</sup> CFU.

<sup>c</sup> Mice living longer than 14 days were assigned a survival time of 14 days for the determination of median values.

## EXAMPLE 12

5 This example described the cloning and expression of a chimeric gene encoding for a chimeric polypeptide corresponding to the carboxy-terminal region of BVH-3 in fusion at the C' end to the carboxy-terminal region of BVH-11 and the additive protection observed after vaccination with a chimeric polypeptide.

10 It is clear from the studies described above that BVH-3 and BVH-11 are serologically distinct molecules simultaneously present on S. pneumoniae. The results of immunological studies of mice indicate that both proteins are good vaccine candidates. These proteins have the potential to provide protection against all pneumococci, regardless of serotype. Even though the two proteins share epitopes and sequences, they have different characteristics and may serve different biological functions. Thus, immunization against the two proteins may provide a higher level of protection than that imparted by each individually. To examine this, several avenues where full-length or truncated BVH-3 and BVH-11 are administered in combination or in conjugation can be explored. Here we describe the 15 genetic engineering of a BVH-3-BVH-11 fusion gene and protein, termed NEW12 (**SEQ ID NO:76** and **SEQ ID NO:58**, respectively), and the potential use of NEW12 protein as a vaccine.

20 **BVH-3** and **BVH-11** gene fragments corresponding to the 3' end of the genes were amplified by PCR using pairs of oligonucleotides engineered to amplify fragments spanning nucleotides 1414 to 3117 (**SEQ ID NO: 1**) and nucleotides 1060 to 2520 (**SEQ ID NO: 3**) from S. pneumoniae strain SP64 **BVH-3** and **BVH-11** genes, respectively. The primers used, HAMJ278 and HAMJ279; HAMJ282 and HAMJ283 had a restriction

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30

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endonuclease site at the 5' end, thereby allowing directional in-frame cloning of the amplified product into the digested pET21b(+) plasmid vector (Table 2). PCR-amplified products were digested with restriction

5 endonucleases and ligated to linearized plasmid pET21b(+) vector digested likewise. The resultant plasmid constructs were confirmed by nucleotide sequence analysis. The recombinant pET21b(+) plasmid containing the NdeI-HindIII BVH-3 PCR product was linearized by digestion with the

10 restriction enzymes HindIII and NotI for the in-frame cloning of the HindIII-NotI DNA fragment obtained from the recombinant pET21(+) vector containing the BVH-11 gene fragment. Clones were first stabilized in E. coli DH5 $\alpha$  before introduction into E. coli BL21(λDE3) for expression

15 of a chimeric pneumococcal protein molecule. The recombinant chimeric polypeptide, termed NEW 12, was expressed as C-terminal fusion with an His-tag. The expressed recombinant NEW 12 protein was purified from supernatant fractions obtained from centrifugation of

20 sonicated IPTG-induced E. coli cultures using a His-Bind metal chelation resin (QIAgen, Chatsworth, CA).

According to the same procedure described above, it is possible to construct other chimeric polypeptides, as a

25 result of a simultaneous expression of New 1 and New 4, New 1 and New 5, New 1 and New 10, or New 1 and New 14. The construction can be with New 1 upstream or downstream of New 4, New 5, New 10, BVH-11B or New 14. It is also possible to construct other chimeric polypeptides as a

30 result of a simultaneous expression of more than two fragments of either genes of BVH-3, BVH-11 or BVH-11-2.

Groups of 8 female BALB/c mice (Charles River) were immunized subcutaneously two times at three-week intervals

35 with 25  $\mu$ g of either affinity purified His-Tag-fusion NEW1,

BVH-11B or NEW12 protein in presence of 15 µg of QuilA adjuvant. Ten to 14 days following the last immunization, the mice were challenged with virulent S. pneumoniae. As demonstrated before, NEW1 and BVH-11B molecules comprising 5 amino acids 472 to 1039 from BVH-3 protein and amino acids 354-840 from BVH-11 protein, respectively, correspond to portions of the proteins capable of eliciting a protective immune response. To determine if a chimeric polypeptide would significantly improve the protection compared with 10 those seen for the individual counterparts, the challenge dose was adjusted in a manner that protection was not expected with NEW1 and BVH-11B molecules. Interestingly, the chimeric NEW12 protein, elicited protection against the mouse-virulent strains WU2 and P4241. Seven out of 8 mice 15 immunized with NEW12 were still alive 10 days after the challenge while 28 out of 32 mice immunized with NEW1, BVH-11B, BVH-3M or adjuvant alone were dead by five days post-challenge. Thus, vaccination of mice with NEW12 provided the highest degree of protection against WU2 challenge. 20 These results indicate that immunization with a chimeric polypeptide and possibly a combination of BVH-3 and BVH-11 gene products can provide additional protection to that obtained by administration of BVH-3 or BVH-11 antigens alone.

25

Table 9. Evaluation of protection elicited by vaccination of mice with the chimeric NEW12 molecule

Immunogen	Challenge with WU2		Challenge with P4241	
	Alive : dead <sup>a</sup>	Median day alive	Alive : dead	Median day alive
None	0 : 8	1	0 : 8	5
NEW1	2 : 6	2	1 : 7	8
BVH-11B	1 : 7	3.5	8 : 0	>14
NEW12	6 : 2	>14	7 : 1	>14
BVH-3M	1 : 7	3	8 : 1	>14

## EXAMPLE 13

5

This example illustrates the identification of additional BVH-3 and BVH-11 related sequences in *Streptococcus* species other than *S. pneumoniae*.

10 It was previously shown that BVH-3, BVH-11 and BVH-11-2 are a family of related proteins sharing common sequences. Homology searches were performed with the nucleotide sequence from the conserved region of these genes and compared with GenBank and EMBL sequences using FASTA. The 15 most significant homology was observed with a 2.469-kb gene coding for a calculated 92-kDa protein (SEQ ID NO: 81) of unknown function in *S. agalactiae* also called group B streptococcus or GBS. The gene was designated BVH-71. A protein demonstrating 99.2% identity and 99.5% similarity 20 with that of GBS was also identified in *S. pyogenes* also called group A streptococcus or GAS (SEQ ID NO: 83). The 5' region of the BVH-71 sequences (SEQ ID NO: 80 and SEQ ID NO: 82), spanning nucleotides 1 to 717, demonstrated 58 and 60% identity with the conserved regions of BVH-3 25 (nucleotides 1 to 675) and BVH-11 (nucleotides 1 to 684) genes respectively. The first 239 amino acids of the translated sequences of the GBS and GAS BVH-71 open reading frames are 51 and 54% identical to the first 225 and 228 amino acids of BVH-3 and BVH-11, respectively. In addition 30 to structural similarities, streptococcal BVH-3, BVH-11 and BVH-71 proteins also share antigenic epitopes. A 97-kDa band was revealed on Western blots of GAS or GBS whole cells, using Mab H11-1.1-G11 reactive with the BVH-3 and BVH-11 conserved regions. Similarly, GAS and GBS

recombinant BVH-71 proteins were detected in Western immunoblot analysis.

These results indicate that BVH-71, BVH-3 and BVH-11 proteins might share similar functions. Our results also 5 suggest that BVH-71 proteins can be used as protein vaccine components of anti-streptococcus. In a further embodiment BVH-71 proteins can be used as protein vaccine components of anti-GAS or anti-GBS vaccines.

What is claimed is:

1. An isolated polynucleotide encoding a polypeptide having at least 70% identity to a second polypeptide having a sequence chosen from: **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.
2. A polynucleotide according to claim 1, wherein said polynucleotide encodes a polypeptide having at least 95% identity to the second polypeptide.
3. An isolated polynucleotide encoding a polypeptide capable of generating antibodies having binding specificity for a polypeptide having a sequence chosen from: **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.
4. An isolated polynucleotide that is complementary to the polynucleotide of claim 1.
5. An isolated polynucleotide that is complementary to the polynucleotide of claim 3.
6. The polynucleotide of claim 1, wherein said polynucleotide is DNA.
7. The polynucleotide of claim 3, wherein said polynucleotide is DNA.
8. The polynucleotide of claim 1, wherein said polynucleotide is RNA.
9. The polynucleotide of claim 3, wherein said polynucleotide is RNA.

10. A vector comprising the polynucleotide of claim 1, wherein said DNA is operably linked to an expression control region.
11. A vector comprising the polynucleotide of claim 3, wherein said DNA is operably linked to an expression control region.
12. A host cell transfected with the vector of claim 10.
13. A host cell transfected with the vector of claim 11.
14. A process for producing a polypeptide comprising culturing a host cell according to claim 12 under conditions suitable for expression of said polypeptide.
15. A process for producing a polypeptide comprising culturing a host cell according to claim 13 under condition suitable for expression of said polypeptide.
16. An isolated polypeptide having at least 70% identity to a second polypeptide having an amino acid sequence chosen from: **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.
17. An isolated polypeptide capable of generating antibodies having binding specificity for a second polypeptide having a sequence chosen from: **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.
18. An isolated polypeptide having an amino acid sequence chosen from **SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83** or fragments, analogs or derivatives thereof.

19. An isolated polypeptide according to claim 18, wherein the N-terminal Met residue is deleted.
20. An isolated polypeptide according to claim 18, wherein the secretory amino acid sequence is deleted.
21. A chimeric polypeptide comprising two or more polypeptides chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.
22. A chimeric polypeptide comprising two or more polypeptides chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof; provided that the polypeptides or fragments, analogs or derivatives thereof are linked as to form a chimeric polypeptide.
23. A chimeric polypeptide of formula (I):  
$$A-(B)_m-(C)_n-D \quad (I)$$
Wherein;  
m is 0 or 1,  
n is 0 or 1,  
A is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;  
B is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof;  
C is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55 to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof; and  
D is chosen from SEQ ID NOS: 2, 4, 6, 8, 10, 14, 16, 55

to 75, 77 to 79, 81, 83 or fragments, analogs or derivatives thereof.

24. A chimeric polypeptide of formula (I):

$A-(B)_m-(C)_n-D$  (I)

Wherein;

$m$  is 0 or 1,

$n$  is 0 or 1,

$A$  is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof;

$B$  is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77, or fragments, analogs or derivatives thereof;

$C$  is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof; and

$D$  is chosen from SEQ ID NOS :10, 58, 60, 62, 64, 67, 68, 69, 72, 74, 77 or fragments, analogs or derivatives thereof.

25. A vaccine composition comprising a polypeptide according to any one of claims 16 to 24 and a pharmaceutically acceptable carrier, diluent or adjuvant.

26. A method for therapeutic or prophylactic treatment of meningitis, otitis media, bacteremia or pneumonia infection in an individual susceptible to meningitis, otitis media, bacteremia or pneumonia infection comprising administering to said individual a therapeutic or prophylactic amount of a composition according to claim 25.

27. A method for therapeutic or prophylactic treatment of streptococcal bacterial infection in an individual

susceptible to streptococcal infection comprising administering to said individual a therapeutic or prophylactic amount of a composition according to claim 25.

28. A method according to claim 26, wherein said individual is a mammal.
29. A method according to claim 27, wherein said individual is a human.
30. A method according to claim 22, wherein said bacterial infection is S.pneumoniae, group A *streptococcus (pyogenes)*, group B *streptococcus* (GBS or *agalactiae*), *dysgalactiae*, *uberis*, *nocardia* or *Staphylococcus aureus*.
31. A method according to claim 26, wherein said bacterial infection is S.pneumoniae.
32. Use of a vaccine composition according to claim 25 for the prophylactic or therapeutic treatment of Streptococcal infection in an animal susceptible to or infected with streptococcal infection comprising administering to said animal a prophylactic or therapeutic amount of the composition.

ATGAAATTAA	GTAAAAAATA	TATAGCAGCT	GGATCAGCTG	TTATCGTATC	CTTGAGTCTA	60
TGTGCCATG	CACTAAACCA	GCATCGTCG	CAGGAAAATA	AGGACAATAA	TCGTGTCTCT	120
TATGTGGATG	GCAGCCAGTC	AAAGTCAGAAA	AGTAAAAC	TGACACCCAGA	CCAGGTTAGC	180
CAGAAAAGAAG	GAATTCAAGGC	TGAGCAAATT	GTAAATCAAAA	TTACAGATCA	GGGCTATGTA	240
ACGTCACACG	GTGACCACTA	TCATTACTAT	AATGGGAAAG	TTCCCTATGA	TGCCCTCTTT	300
AGTGAAGAAC	TCTTGATGAA	GGATCCAAC	TATCAACTTA	AAGACGCTGA	TATTGTCAAT	360
GAAGTCAGG	GTGGTTATAT	CATCAAGGTC	GATGGAAAAT	ATTATGTCTA	CCTGAAAGAT	420
GCAGCTCATG	CTGATAATGT	TCGAACTAA	GATGAAATCA	ATCGTCAAAA	ACAAGAACAT	480
GTCAAAGATA	ATGAGAAGGT	TAACTCTAAT	GTTGCTGTAG	CAAGGTCTCA	GGGACGATAT	540
ACGACAAATG	ATGGTTATGT	CTTTAATCCA	GCTGATATTAA	TCGAAGATAC	GGGTAAATGCT	600
TATATCGTTC	CTCATGGAGG	TCACTATCAC	TACATTCCCA	AAAGCGATT	ATCTGCTAGT	660
GAATTAGCAG	CAGCTAAAGC	ACATCTGGCT	GGAAAAAAATA	TGCAACCGAG	TCAGTTAACG	720
TATTCTCAA	CAGCTAGTGA	CAATAACACG	CAATCTGTAG	CAAAGGATC	AACTAGCAAG	780
CCAGCAAATA	AATCTGAAA	TCTCCAGAGT	CTTTGAAGG	AACTCTATGA	TTCACCTAGC	840
GCCCAACGTT	ACAGTGAATC	AGATGCCCTG	GTCTTGACCC	CTGCTAAGAT	TATCAGTCGT	900
ACACCAAATG	GAGTTGCGAT	TCGGCATGGC	GACCATTACC	ACTTTATTCC	TTACAGCAAG	960
CTTTCTGCTT	TAGAAGAAA	GATTGCCAGA	ATGGTGCCTA	TCAGTGGAAC	TGGTTCTACA	1020
GTTCCTACAA	ATGCAAAACC	TAATGAAGTA	GTGTCTAGTC	TAGGCAGTCT	TTCAAGCAAT	1080
CTTTCTCTT	TAACGACAAG	TAAGGAGCTC	TCTTCAGCAT	CTGATGGTTA	TATTTTTAAT	1140
CCAAAAGATA	TCGTTGAAGA	AACGGCTACA	GCTTATATTG	TAAGACATGG	TGATCATTTTC	1200
CATTACATT	CAAATCAAA	TCAAATTGGG	CAACCGACTC	TTCCAAACAA	TAGTCTAGCA	1260
ACACCTTCTC	CATCTCTTCC	AATCAATCCA	GGAACTTCAC	ATGAGAAACA	TGAAGAAGAT	1320
GGATACGGAT	TTGATGCTAA	TCGTATTATC	GCTGAAAGATG	AATCAGGTT	TGTCATGAGT	1380
CACGGAGACC	ACAATCATTA	TTTCTTCAGA	AAGGACTTGA	CAGAAGAGCA	AAATTAAAGGCT	1440
GCGCAAAAC	ATTTAGAGGA	AGTTAAAAC	AGTCATAATG	GATTAGATTC	TTTGTCATCT	1500
CATGAACAGG	ATTATCCAGG	TAATGCCAA	GAATGAAAG	ATTTAGATAA	AAAATCGAA	1560
GAAAAAATTG	CTGGCATTAT	GAAACAAATAT	GGTGTCAAAC	GTGAAAGTAT	TGTCGTGAAT	1620
AAAGAAAAAA	ATGCGATTAT	TTATCCGCAT	GGAGATCACC	ATCATGCAGA	TCCGATTGAT	1680
GAACATAAAC	CGGTTGGAAT	TGGTCATTCT	CACAGTAACT	ATGAACTGTT	TAAACCCGAA	1740
GAAGGAGTTG	CTAAAAAAGA	AGGGATAAA	GTTTATACG	GAGAAGAATT	AACGAATGTT	1800
GTAAATTGT	AAAAAAATAG	TACGTTAA	AATCAAAAC	TTACTCTAGC	CAATGGTCAA	1860
AAACCGCTTT	CTTTTAGTTT	TCGGCCTGAA	TTGGAGAAAA	AATTAGGTAT	CAATATGCTA	1920
GTAAAATTAA	TAACACCAGA	TGGAAAAGTA	TTGGAGAAAG	TATCTGGTAA	AGTATTTGGA	1980
GAAGGAGTAG	GGAATATTGC	AAACTTTGAA	TTAGATCAAC	CTTATTAC	AGGACAAACA	2040
TTTAAGTATA	CTATCGCTTC	AAAAGATTAT	CCAGAAGTAA	GTTATGATGG	TACATTACAA	2100
GTTCCAAACCT	CTTTAGCTTA	AAAATGGCC	AGTCAAACGA	TTTCTATCC	TTTCCATGCA	2160
GGGGATACTT	ATTTAAGAGT	GAACCCCTAA	TTTGCAGTGC	CTAAAGGAAC	TGATGCTTTA	2220
GTCAGAGTGT	TTGATGAATT	TCATGGAAAT	GCTTATTTAG	AAAATAACTA	TAAGTTGGT	2280
GAAATCAAAT	TACCGATTCC	GAAATTAAAC	CAAGGAACAA	CCAGAACGGC	CGGAAATAAA	2340
ATTCCGTAA	CCTTCATGGC	AAATGCTTAT	TTGGACAATC	AATCGACTTA	TATTGTGGAA	2400
GTACCTATCT	TGGAAAAAAGA	AAATCAAAC	GATAAACCAA	GTATTCTACC	ACAATTAA	2460
AGGAATAAAAG	CACAAGAAAA	CTCAAAACTT	GATGAAAAGG	TAGAAGAAC	AAAGACTAGT	2520
GAGAAGGTAG	AAAAAGAAAA	ACTTTCTGAA	ACTGGGAATA	GTACTAGTAA	TTCAACGTTA	2580
GAAGAAGTTC	CTACAGTGG	TCCTGTACAA	GAAAAAGTAG	CAAATTGTC	TGAAAGTTAT	2640
GGGATGAAGC	TAGAAAATGT	CTTGTAAAT	ATGGACGGAA	CAATTGAATT	ATATTACCA	2700
TCAGGAGAAG	TCATTTAAA	GAATATGGCA	GATTTACAG	GAGAAGCACC	TCAAGGAAAT	2760
GGTGAATAA	AACCATCTGA	AAATGGAAAA	GTATCTACTG	GAACAGTGA	GAACCAACCA	2820
ACAGAAAATA	AACCAGCAGA	TTCTTACCA	GAGGCACCAA	ACGAAAAC	TGTAAGAACCA	2880
GAAAACCTAA	CGGATAATGG	AATGTTGAAT	CCAGAAGGGG	ATGTGGGGAG	TGACCCCTATG	2940
TTAGATCCAG	CATTAGAGGA	AGCTCCAGCA	GTAGATCCTG	TACAAGAAA	ATTAGAAAAA	3000
TTTACAGCTA	GTTACGGATT	AGGCTTAGAT	AGTGTATAT	TCAATATGGA	TGGAACCGATT	3060
GAATTAAGAT	TGCCAAGTGG	AGAAGTGATA	AAAAAGAATT	TATCTGATT	CATAGCGTAA	3120

(SEQ ID NO: 1)

FIGURE 1

MKFSKKYIAA	GSAVIVSLSL	CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	50
SENLTPDQVS	QKEGIQAEQI	VIKITDQGYV	TSHGDHYHY	NGKVPYDALF	100
SEELLMKDPN	YQLKDADIVN	EVKGYYIIV	DGKYYVYLKD	AAHADNVRTK	150
DEINRQKQEH	VKDNEKVNSN	VAVARSQGRY	TTNDGYVFNP	ADIIIEDTGNA	200
YIVPHGGHYH	YIPKSDLSAS	ELAAAKAHLA	GKNMQPSQLS	YSSTASDNNT	250
QSVAKGSTSK	PANKSENLQS	LLKELYDSPS	AQRYSESDSL	VFDPAKIISR	300
TPNGVAIPHG	DHYHFIPYSK	LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	350
VSSLGSLSSN	PSSLTTSKEL	SSASDGYIFN	PKDIVEETAT	AYIVRHDHF	400
HYIPKSNQIG	QPTLPNNSLA	TPSPSLPINP	GTSHEKHEED	GYGFDANRII	450
AEDESGFVMS	HGDHNHYFFK	KDLTEEQIKA	AQKHLEEVKT	SHNGLDSLSS	500
HEQDYPGNAK	EMKDLDKKIE	EKIAGIMKQY	GVKRESIVVN	KEKNATIYPH	550
GDHHADPID	EHKPVGIGHS	HSNYELFKPE	EGVAKKEGNK	VYTGEELTNV	600
VNLLKNSTFN	NQNFTLNGQ	KRVSFSFPPE	LEKKLGINML	VKLITPDGKV	650
LEKVSGKVFG	EGVGNIANFE	LDQPYLPGQT	FKYTIASKDY	PEVSYDGTFT	700
VPTSLAYKMA	SQTIFYPFHA	GDTYLRVNPO	FAVPKGTDAL	VRVFDEFHGN	750
AYLENNYKVG	EIKLPIPKLN	QGTTRTAGNK	IPVTFMANAY	LDNQSTYIVE	800
VPILEKENQT	DKPSILPQFK	RNKAQENSKL	DEKVEEPKTS	EKVEKEKLSE	850
TGNSTSNSTL	EEVPTVDPVQ	EKVAKFAESY	GMKLENVLFN	MDGTIELYLP	900
SGEVIKKNMA	DFTGEAPQGN	GENKSENGK	VSTGTVENQP	TENKPADSLP	950
EAPNEKPVKP	ENSTDNGMLN	PEGNVGSDPM	LDPALEEAPA	VDPVQEKLEK	1000
FTASYGLGLD	SVIFNMDGTI	ELRLPSGEVI	KKNLSDFIA	(SEQ ID NO: 2)	1039

FIGURE 2

ATGAAAATCA	ATAAAAATA	TCTAGCTGGG	TCAGTAGCTA	CACTTGT	TTT	AAGTGTCTGT	60
GCTTATGAAC	TAGGTTGCA	TCAAGCTCAA	ACTGTAAAAG	AAAATAATCG	TGTTTCCTAT	120	
ATAGATGGAA	ACAAAGCGAC	GCAAAAAACG	GAGAATTGAA	CTCCGTATGA	GGTTAGCAAG	180	
CGTGAAGGAA	TCAACGCCGA	ACAAATCGTC	ATCAAGATTA	CGGATCAAGG	TTATGTGACC	240	
TCTCATGGAG	ACCATTATCA	TTACTATAAT	GGCAAGGTCC	CTTATGTGTC	CATCATCAGT	300	
GAAGAGCTCC	TCATGAAAGA	TCCGAATTAT	CAGTTGAAGG	ATTCAGACAT	TGTCAATGAA	360	
ATCAAGGGTG	GTTATGTCTAT	TAAGTAAAC	GGTAAATACT	ATGTTTACCT	TAAGGATGCA	420	
GCTCATGCCG	ATAATGTCG	TACAAAAGAA	GAAATCAATC	GGCAAAAACA	AGAACATAGT	480	
CAGCATCGTG	AAGGAGGGAC	TTCAGCAAAC	GATGGTGC	TAGCCTTGC	ACGTTCACAG	540	
GGACGCTACA	CCACAGATGA	TGGTTATATC	TTCAATGCAT	CTGATATCAT	CGAAGATAACG	600	
GGCGATGCCT	ATATCGTCC	TCATGGAGAT	CATTACCAT	ACATTCTAA	GAATGAGTTA	660	
TCAGCTAGCG	AGTTGGCTGC	TGCAAGGCC	TTCCCTATCTG	GTCGGGAAA	TCTGTCAAAT	720	
TTAAGAACCT	ATCGCCGACA	AAATAGCGAT	AAACACTCCAA	GAACAAACTG	GGTACCTCT	780	
GTAAGCAATC	CAGGAACTAC	AAATACAAAC	ACAAGCAACA	ACAGCAACAC	TAACAGTCAA	840	
GCAAGTCAAA	GTAATGACAT	TGATAGTCTC	TTGAAAACAGC	TCTACAAACT	GCCTTTGAGT	900	
CAACGCCATG	TAGAATCTGA	TGGCCTTATT	TTCGACCCAG	CGCAAATCAC	AAGTCGAACC	960	
GCCAGAGGTG	TAGCTGTCCC	TCATGGTAAC	CATTACCAT	TTATCCCTTA	TGAACAAATG	1020	
TCTGAATTGG	AAAAACGAAT	TGCTCGTATT	ATTCCCCTTC	GTATCGTTC	AAACCATTGG	1080	
GTACCAAGATT	CAAGACCAGA	AGAACCAAGT	CCACAAACCGA	CTCCAGAAC	TAGTCCAAGT	1140	
CCGCAACCTG	CACCAAATCC	TCAACCAGCT	CCAAGCAATC	CAATTGATGA	GAATTGGTC	1200	
AAAGAAGCTG	TTCGAAAAGT	AGGCATGGT	TATGTCTTG	AGGAGAATGG	AGTTTCTCGT	1260	
TATATCCCAG	CCAAGAATCT	TTCAGCAGAA	ACAGCAGCAG	GCATTGATAG	CAAACGGCC	1320	
AAGCAGGAAA	GTTTATCTCA	TAAGCTAGGA	GCTAAGAAA	CTGACCTCCC	ATCTAGTGT	1380	
CGAGAATTTC	ACAATAAGGC	TTATGACTTA	CTAGCAAGAA	TTCACCAAGA	TTTACTTGAT	1440	
AATAAAAGGTC	GACAAGTTGA	TTTGAGGCT	TTGGATAACC	TGTTGGAACG	ACTCAAGGAT	1500	
GTCTCAAGTG	ATAAAAGTCAA	GTTAGTGGAT	GATATTCTTG	CCTTCTTAGC	TCCGATTCGT	1560	
CATCCAGAAC	GTTTAGGAAA	ACCAAATGCG	CAAATTACCT	ACACTGATGA	TGAGATTCAA	1620	
GTAGCCAAGT	TGGCAGGCAA	GTACACAAAC	GAAGACGGTT	ATATCTTG	TCCTCGTGT	1680	
ATAACCAGTG	ATGAGGGGGA	TGCCTATGTA	ACTCCACATA	TGACCCATAG	CCACTGGATT	1740	
AAAAAAAGATA	GTTTGTCTGA	AGCTGAGAGA	GCGGCAGCCC	AGGTTATGC	AAAGAGAAA	1800	
GGTTTGACCC	CTCCCTCGAC	AGACCATCAG	GATTCAAGGAA	ATACTGAGGC	AAAAGGAGCA	1860	
GAAGCTATCT	ACAACCGCGT	GAAAGCAGCT	AAGAAGGTGC	CACTTGATCG	TATGCCCTAC	1920	
AATCTTCAAT	ATACTGTAGA	AGTCAAAAC	GGTAGTTAA	TCATACCTCA	TTATGACCAT	1980	
TACCATATAAC	TCAAATTTGA	GTGGTTTGAC	GAAGGCCTTT	ATGAGGCACC	TAAGGGGTAT	2040	
ACTCTTGAGG	ATCTTTGGC	GACTGTCAAG	TACTATGTG	AACATCCAAA	CGAACGTCCG	2100	
CATTAGATA	ATGGTTTG	TAACGCTAGC	GACCATGTT	AAAGAAACAA	AAATGGTCAA	2160	
GCTGATACCA	ATCAAACGGA	AAAACCAAGC	GAGGAGAAC	CTCAGACAGA	AAAACCTGAG	2220	
GAAGAAACCC	CTCGAGAAGA	GAAACCACAA	AGCGAGAAC	CAGAGTCTCC	AAAACCAACA	2280	
GAGGAACCAG	AAGAAGAAC	ACCAGAGGAA	TCAGAAGAAC	CTCAGGTCGA	GACTGAAAAG	2340	
GTTGAAGAAA	AACTGAGAGA	GGCTGAAGAT	TTACTTGGAA	AAATCCAGGA	TCCAATTATC	2400	
AAGTCCAATG	CCAAAGAGAC	TCTCACAGGA	TTAAAAAATA	ATTTACTATT	TGGCACCCAG	2460	
GACAACAATA	CTATTATGGC	AGAAGCTGAA	AAACTATTGG	CTTTATTAAA	GGAGAGTAAG	2520	
TAA	(SEQ ID NO: 3)					2523	

FIGURE 3

MKINKKYL	AG SVATLVL SVC	AYELGLHQAQ	TVKENN RV SY	IDGKQATQKT	50
ENLTPDEV SK	REGINA EQIV	IKITDQGYVT	SHGDHYH YN	GKVPYDAI IS	100
BELLMKD PNY	QLKDSDIV NE	IKGGYVIKVN	GKYYVYL KDA	AHADNVRTKE	150
EINRQKQEH S	QHREGGTSAN	DGAVAFARSQ	GRYTTDDGYI	FNASDIIEDT	200
GDAYIVPHGD	HYHYIPKNE L	SASELAAA EA	FLSGRENLSN	LRTYRRQNSD	250
NTPRTNWVPS	VSNPGTTNTN	TSNNNSNTNSQ	ASQSNDIDSL	LKQLYKLPLS	300
QRHVESDGLI	FDPAQITSRT	ARGVAVPHGN	HYHFIPYEQM	SELEKRIARI	350
IPLRYRSNH W	VPDSRPEEPS	PQPTPEPSPS	PQPAPNPQPA	PSNPIDEKLV	400
KEAVRKVG DQ	YVFEENGVSR	YIPAKNLSAE	TAAGIDS KLA	KQESLSHKLG	450
AKKTDL PSSD	REFYNKAYDL	LARIHQD LLD	NKGRQVD FEA	LDNLLE RLKD	500
VSSDKV KLVD	DILAFLAPI R	HPERLGK PNA	QITYTDDEI Q	VAKLAGKYTT	550
EDGYIFD PRD	ITSDEGDAY V	TPHMTHSHWI	KKDSLSE AER	AAAQAYAKEK	600
GLTPPSTDHQ	DSGNTEAKGA	EAIYNRVKA	KKVPLDRMPY	NLQYTVBVKN	650
GSLIIPH YDH	YHNIKFEW FD	EGLYEAPK GY	TLEDLLATV K	YYVEHPN ERP	700
HSDNGFGN AS	DHVQRNKNG Q	ADTNQTEKPS	EEKPKQTEKPE	EETPREEK PQ	750
SEKPESPKPT	EEPEEE SPEE	SEEPQVETEK	VEEKLREAED	LLGKIQDPII	800
KSNAKETLTG	LKNNNLLFGT Q	DNNTIMAEAE	KLLALLKESK	(SEQ ID NO: 4)	840

FIGURE 4

ATGGAGAATA	TAGACATGTT	AAAATCAAAT	CATGAGCGAA	GAATGCCGTTA	TTCCATT CGT	60
AAATTAGTG	TAGGAGTAGC	TAGCGTAGCT	GTTGCCAGTC	TTTTTATGGG	AA GTGTTGTA	120
CATGCGACAG	AGAAAAGAGGG	AA GTACCCAA	GCAGCCACTT	CTTTTAATAG	GGGAAATGGA	180
AGTCAGG CAG	ACAAACGTGG	AGAAACTCGAT	TTAGAACGAG	ATAAGGCAAT	GAAAGCGGTC	240
AGTGAATATG	TAGGAAAAT	GGTGAGAGAT	GCCTATGTAA	AATCAGATAG	AAAAGGACAT	300
AAAATACTG	TAGCTCTAGT	TAACCAGTTG	GGAAACATTA	AGAACAGGT A	TTTGAATGAA	360
ATAGTT CATT	CAACCTCAAA	AAGCCA ACTA	CAGGA ACTG A	TGATGAAGAG	TCAATCAGAA	420
GTAGATGAAG	CTGTGTCTAA	ATTTGAAAAG	GACTCATT TT	CTTCGTCAAG	TTCAGGATCC	480
TCCACTAAAC	CAGAAACTCC	GCAGCCGGAA	AATCCAGAGC	ATCAAA AAC	ACAAACTCCA	540
TCTCCGGATA	CCAAACCAAG	CCCTCAACCA	GAAGGCAAGA	AACCAAGCGT	ACCAGACATT	600
AATCAGGAAA	AAGAAAAGC	TAAGCTTGCT	GTAGTAACCT	ACATGAGCAA	GATTTTAGAT	660
GATATACAAA	AACATCATCT	GCAGAAAGAA	AAACATCGTC	AGATTGTTGC	TCTTATTAAG	720
GAGCTTGATG	AGCTTAAAAA	GCAAGCTCTT	TCTGAAATTG	ATAATG TAA	TACCAAAGTA	780
GAAATTGAAA	ATACAGTCCA	CAAGATATT	GCAGACATGG	ATGCAGTTGT	GACTAAATTC	840
AAAAAAGGCT	TAACTCAGGA	CACACCAAA	GAACCAGGT A	ACAAAAAAC	ATCTGCTCCA	900
AAACCAGGTA	TGCAACCAAG	TCCTCAACCA	GAGGTAAAC	CGCAGCTGGA	AAAACCAAA	960
CCAGAGGTTA	AACCGCAACC	AGAAAAACCA	AAACCAGAGG	TTAAACCGCA	GCCGGAAAAA	1020
CCAAAACCA G	AGGTTAAACC	GCAGCCGGAA	AAACCAAAAC	CAGAGGTTAA	ACCGCAGCCG	1080
GAAAAAACCA A	AACCAGAGGT	TAACCCGCA G	CCGGAAAAAC	CAAAACCAGA	GGTTAAACCG	1140
CAGCCGGAAA	AACCAAAACC	AGAGGTTAAA	CCGCAGCCGG	AAAAACCAA	ACCAAGAGGT	1200
AAACCGCAGC	CGGAAAAACCA	AAAACCAAGAG	GTAAACCGC	AGCCGAAAA	ACCAAAACCA	1260
GAGGTTAAAC	CGCAGCCGG A	AAAACCAA A	CCAGAGGTT A	AACCGCAACC	AGAAAAACCA	1320
AAACCAGAGG	TTAAACCGCA	ACCAGAAAAA	CCAAAACCA G	ATAATAGCAA	GCCACAAGCA	1380
GATGATAAGA	AGCCATCAAC	TACAATAAT	TTAAGCAAGG	ACAAGCAACC	TTCTAACCAA	1440
GCTTCAACAA	ACGAAAAGC	AACAAATAA	CCGAAGAAGT	CATTGCCATC	AACTGGATCT	1500
ATTTCAAATC	TAGCACTTGA	AATTGCAAGGT	CTTCTTACCT	TGGCGGGGGC	AACCATTCTT	1560
GCTAAGAAA	GAATGAAATA	G	(SEQ ID NO: 5)			1581

FIGURE 5

MENIDMFKSN	HERRMRYSIR	KFSVGVASVA	VASLFMGSVV	HATEKEGSTQ	50
AATSFNRGNG	SQAEQRGELD	LERDKAMKAV	SEYVGKMRD	AYVKSDRKRH	100
KNTVALVNQL	GNIKNRYLNE	IVHSTSCKSQL	QELMMKSQSE	VDEAVSKFEK	150
DSFSSSSSGS	STKPETPQPE	NPEHQKPTTP	SPDTKPSQQP	EGKKPSVPDI	200
NQEKEKAKLA	VVTYMSKILD	DIQKHHLQKE	KHRQIVALIK	ELDELKKQAL	250
SEIDNVNTKV	EIENTVHKIF	ADMADAVTKF	KKGLTQDTPK	EPGNKKPSAP	300
KPGMQPSPQP	BVKPQLEKPK	PEVKPQPEKP	KPEVKPQPEK	PKPEVKPQPE	350
KPKPEVKPQP	EKPKPEVKPQ	PEKPKPEVK	QPEKPKPEV	PQPEKPKPEV	400
KPQPEKPKPE	VVKPQPEKPK	EVVKPQPEKPK	PEVKPQPEK	KPEVKPQPEK	450
PKPDNSKPQA	DDKKPSTTNN	LSKDKQPSNQ	ASTNEKATNK	PKKSLPSTGS	500
ISNLALEIAG	LLTAGATIL	AKKRMK	(SEQ ID NO: 6)		526

FIGURE 6

ATGAAATTTA	GTAAAAAATA	TATAGCAGCT	GGATCAGCTG	TTATCGTATC	CTTGAGTCTA	60
TGTGCCTATG	CACTAAACCA	GCATCGTTCG	CAGGAAAATA	AGGACAATAA	TCGTGTCTCT	120
TATGTGGATG	GCAGCCAGTC	AAAGTCAGAAA	AGTAAAAC	TGACACCCAGA	CCAGGTTAGC	180
CAGAAAAGAAG	GAATTTCAGGC	TGAGCAAATT	GTAATCAAAA	TTACAGATCA	GGGCTATGTA	240
ACGTACACACG	GTGACCACTA	TCATTACTAT	AATGGGAAAG	TTCCCTTATGA	TGCCCTCTTT	300
AGTGAAGAAC	TCTTGATGAA	GGATCCAACAC	TATCAACTTA	AAGACGCTGA	TATTGTCAAT	360
GAAGTCAGG	GTGGTTATAT	CATCAAGGTC	GATGGAAAAT	ATTATGCTA	CTTGAAAGAT	420
GCAGCTCATG	CTGATAATG	TCGAACCTAA	GATGAAATCA	ATCGTCAAAA	ACAAGAACAT	480
GTCAAAGATA	ATGAGAAGGT	TAACTCTAA	GTGCTGTAG	CAAGGCTCTCA	GGGACGATAT	540
ACGACAAATG	ATGGTTATG	CTTTAATCCA	GCTGATATTAA	TCGAAGATAC	GGGTAATGCT	600
TATATCGTTC	CTCATGGAGG	TCACTATCAC	TACATTCCCA	AAAGCGATT	ATCTGCTAGT	660
GAATTAGCAG	CAGCTAAAGC	ACATCTGGCT	GGAAAAAATA	TGCAACCGAG	TCAGTTAACG	720
TATTCTCAA	CAGCTAGTGA	CAATAACACG	CAATCTGTAG	CAAAGGATC	AACTAGCAAG	780
CCAGCAAATA	AATCTGAAAA	TCTCCAGAGT	CTTTGAAGG	AACTCTATGA	TTCACCTAGC	840
GCCCCAACGTT	ACAGTGAATC	AGATGGCTG	GTCCTTGACC	CTGCTAAGAT	TATCAGTCGT	900
ACACCAAATG	GAGTTGCGAT	TCCGCATGGC	GACCATTACC	ACTTTATTCC	TTACAGCAAG	960
CTTTCTGCTT	TAGAAGAAAA	GATTGCCAGA	ATGGTGCCTA	TCAGTGAAC	TGGTTCTACA	1020
GTTTCTACAA	ATGCAAAACC	TAATGAAGTA	GTGTCTAGTC	TAGGCAGTCT	TTCAAGCAAT	1080
CCTTCTTCTT	TAACGACAAG	TAAGGAGCTC	TCTTCAGCAT	CTGATGGTTA	TATTTTTAAT	1140
CCAAAAGATA	TCGTTGAAGA	AACGGCTACA	GCTTATATTG	TAAGACATGG	TGATCATTTC	1200
CATTACATTC	CAAATCAAA	TCAAATTGGG	CAACCGACTC	TTCCAAACAA	TAGTCTAGCA	1260
ACACCTCTC	CATCTCTTCC	AATCAATCCA	GGAACCTTCAC	ATGAGAAACA	TGAAGAAGAT	1320
GGATACGGAT	TTGATGCTAA	TCGTATTATC	GCTGAAGATG	AATCAGGTTT	TGTCATGAGT	1380
CACGGAGACC	ACAATCATTA	TTTCTTCAG	AAGGACTTGA	CAGAAGAGCA	AATTAAGGTG	1440
CGAAAAAACAA	TTTAG	(SEQ ID NO: 7)			1455	

FIGURE 7

MKFSKKYIAA GSAVIVSLSL CAYALNQHRS QENKDNNRVS YVDGSQSSQK	50
SENLTQDQVS QKEGIQAEQI VIKITDQGYV TSHGDHYHYY NGKVPYDALF	100
SEELLMKDPN YQLKDADIVN EVKGGYIIKV DGKYYVYLKD AAHADNVRTK	150
DEINRQKQEH VKDNEKVNSN VAVARSQGRY TTNDGYVFNP ADIIIEDTGN	200
YIVPHGGHYH YIPKSDLSAS ELAAAKAHLA GKNMQPSQLS YSSTASDNNT	250
QSVAKGSTSK PANKSENLQS LLKELYDSPS AQRYSESDGL VFDPAKIISR	300
TPNGVAIPHG DHYHFIPYSK LSALEEKIAR MVPISGTGST VSTNAKPNEV	350
VSSLGSLSSN PSSLTTSKEL SSASDGYIFN PKDIVEETAT AYIVRHGDHF	400
HYIPKSNQIG QPTLPNNSLA TPSPLPINP GTSHEKHEED GYGF DANRII	450
AEDESGFVMS HGDHNHYFFK KDLTEEQIKV RKNI (SEQ ID NO: 8)	484

FIGURE 8

ATGAAAGATT TAGATAAAAA AATCGAAGAA AAAATTGCTG GCATTATGAA ACAATATGGT	60
GTCAAACGTG AAAGTATTGT CGTGAATAAA GAAAAAAATG CGATTATTAA TCCGCATGGA	120
GATCACCATC ATGCAGATCC GATTGATGAA CATAAACCGG TTGGAATTGG TCATTCTCAC	180
AGTAATCTATG AACTGTTAA ACCCGAAGAA GGAGTTGCTA AAAAAGAAGG GAATAAAAGTT	240
TATACTGGAG AAGAATTAAAC GAATGTTGTT AATTGTTAA AAAATAGTAC GTTTAATAAT	300
CAAAACTTTA CTCTAGCCAA TGGTCAAAAA CGCGTTCTT TTAGTTTCC GCCTGAATTG	360
GAGAAAAAAAT TAGGTATCAA TATGCTAGTA AAATTAATAA CACAGATGG AAAAGTATTG	420
GAGAAAGTAT CTGGTAAAGT ATTTGGAGAA GGAGTAGGGG ATATTGCAAA CTTTGAATTA	480
GATCAACCTT ATTTACCAAGG ACAAAACATT AAGTATACTA TCGCTCAAA AGATTATCCA	540
GAAGTAAGTT ATGATGGTAC ATTTACAGTT CCAACCTTT TAGCTTACAA AATGGCCAGT	600
CAAACGATTT TCTATCCTT CCATGCAGGG GATACTTATT TAAGAGTGAA CCCTCAATT	660
GCAGTGCCTA AAGGAACGTG TGCTTAGTC AGAGTGTGTT ATGAATTTCAT TGAAATGCT	720
TATTTAGAAA ATAACATATAA AGTGTGTTGAA ATCAAATTAC CGATTCCGAA ATTAAACCAA	780
GGAACACCA GAACGGCCGG AAATAAAATT CCTGTAACCT TCATGGCAAA TGCTTATTTG	840
GACAATCAAT CGACTTATAT TGTGGAAGTA CCTATCTTG AAAAGAAAA TCAAACGTAT	900
AAACCAAGTA TTCTACCACA ATTTAAAAGG AATAAACGAC AAGAAAACTC AAAACTTGAT	960
GAAAAGGTAG AAGAACACAA GACTAGTGAG AAGGTAGAAA AAGAAAAACT TTCTGAAACT	1020
GGGAATAGTA CTAGTAATTCA AACGTTAGAA GAAGTTCCTA CAGTGGATCC TGACAAGAA	1080
AAAGTAGCAA AATTGCTGA AAGTTATGGG ATGAAGCTAG AAAATGTCTT GTTTAATATG	1140
GACGGAACAA TTGAATTATA TTACCATCA GGAGAAGTCA TTAAAAAGAA TATGGCAGAT	1200
TTTACAGGAG AAGCACCTCA AGGAAATGGT GAAAATAAAC CATCTGAAAA TGAAAAGTA	1260
TCTACTGGAA CAGTTGAGAA CCAACCAACA GAAAATAAAC CAGCAGATTTC TTTACCAGAG	1320
GCACCAAACG AAAAACCTGT AAAACCAAGAA AACTCAACGG ATAATGGAAT GTTGAATCCA	1380
GAAGGGAATG TGGGGAGTGA CCCTATGTTA GATCCAGCAT TAGAGGAAGC TCCAGCAGTA	1440
GATCCTGTAC AAGAAAAATT AGAAAAATT ACAGCTAGT ACGGATTAGG CTTAGATAGT	1500
GTTATATTCA ATATGGATGG AACGATTGAA TTAAGATTGC CAAGTGGAGA AGTGTATAAAA	1560
AAGAATTATCTGATTTCA AGCGTAA (SEQ ID NO: 9)	1587

FIGURE 9

MKDLDKKIEE	KIAGIMKQYG	VKRESIVVNK	EKNAIYYPHG	DHHADPIDE	50
HKPVGIGHSH	SNYELFKPEE	GVAKKEGNKV	YTGEELTNVV	NLLKNSTFNN	100
QNFTLANGQK	RVSFSFPPEL	EKKLGINMLV	KLITPDGKVL	EKVSGKVFGE	150
GVGNIANFEL	DQPYLPQTF	KYTIASKDYP	EVSYDGTFTV	PTSLAYKMAS	200
QTIFYPFHAG	DTYLRVNPQF	AVPKGTDALV	RVFDEFHGNA	YLENNYKVGE	250
IKLPIPKLNQ	GTTRTAGNKI	PVTFMANAYL	DNQSTYIVEV	PILEKENQTD	300
KPSILPQFKR	NKAQENSKLD	EKVEEPKTSE	KVEKEKLSET	GNSTSNSTLE	350
EVPTVDPVQE	KVAKFAESYG	MKLENVLFN	DGTIELYLPS	GEVIKKNMAD	400
FTGEAPQGNG	ENKPSENGKV	STGTVENQPT	ENKPADSLPE	APNEKPVKPE	450
NSTDNGMLNP	EGNVGSDPML	DPALEEAPAV	DPVQEKLEKF	TASYGLGLDS	500
VIFNMDGTIE	LRLPSGEVIK	KNLSDFIA	(SEQ ID NO: 10)		528

FIGURE 10

BVH3 WU2	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3 RX1	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3 JNR7/87	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3 SP64	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3 P4241	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
BVH3 A66	1	CAYALNQHRSQENKDNNRVSYVDGSSQKSENLT	PDQVSQKEGIQAEQIVIKITDQGYV	60
	*****			*****
BVH3 WU2	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
BVH3 RX1	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
BVH3 JNR7/87	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
BVH3 SP64	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
BVH3 P4241	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
BVH3 A66	61	TSHGDHYHYYNGKVPYDALFSEELLMDPNYQLK	DADIVNEVKGGYIIVKVDGKYYVYLN	120
	*****			*****
BVH3 WU2	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
BVH3 RX1	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
BVH3 JNR7/87	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
BVH3 SP64	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
BVH3 P4241	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
BVH3 A66	121	AAHADNVRTKDEINRQKQEHVKDNEKVNSNA	VARSQGRYTTNDGYVFPNPADIIEDTGNA	180
	*****			*****
BVH3 WU2	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
BVH3 RX1	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
BVH3 JNR7/87	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
BVH3 SP64	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
BVH3 P4241	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
BVH3 A66	181	YIVPHRGHGHYIIPKSDLSASELAAAKAHLAG	KNMQPSQLSYSSTASDNNNTQSVAKGTSK	240
	*****			*****
BVH3 WU2	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3 RX1	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3 JNR7/87	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3 SP64	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3 P4241	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
BVH3 A66	241	PANKSENLQSLLKELYDPSAQRYS	ESDGLVFPDPAKIISRTPNGVAIPHGDHYHFIPYSK	300
	*****			*****
BVH3 WU2	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
BVH3 RX1	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
BVH3 JNR7/87	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
BVH3 SP64	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
BVH3 P4241	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
BVH3 A66	301	LSEALEEKIARMVPISGTGSTVSTNAKPNEV	VSSLGSLSNNPSSLTTSKELSSASDGYIFN	360
	*****			*****
BVH3 WU2	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
BVH3 RX1	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
BVH3 JNR7/87	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
BVH3 SP64	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
BVH3 P4241	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
BVH3 A66	361	PKDIVEETATAYIVRHGDH	PHYIPKSNQIGQPTLPNNSLATPSPLPINPGTSHEKHEED	420
	*****			*****
BVH3 WU2	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
BVH3 RX1	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
BVH3 JNR7/87	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
BVH3 SP64	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
BVH3 P4241	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
BVH3 A66	421	GYGFDANRIIAEDESGFVMSHG	DHNHYFFKKDLTEEQIKAAQKHL	480
	*****			*****

BVH3 WU2	481	HEQDYPNSAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
BVH3 RX1	481	HEQDYPGNAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
BVH3 JNR7/87	481	HEQDYPNSAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
BVH3 SP64	481	HEQDYPGNAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
BVH3 P4241	481	HEQDYPNSAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
BVH3 A66	481	HEQDYPNSAKEMDLDKK1EEK1AGIMKQYGVKRESIVVNEKNAIIYPHGDHHHADPID	540
*****			
BVH3 WU2	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
BVH3 RX1	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
BVH3 JNR7/87	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
BVH3 SP64	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
BVH3 P4241	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
BVH3 A66	541	EHKPGVIGHSHSNYELFKPEEGVAKKEGNKVTGEELTNVNVNLLKNSTFNNQNFTLANGQ	600
*****			
BVH3 WU2	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
BVH3 RX1	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
BVH3 JNR7/87	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
BVH3 SP64	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
BVH3 P4241	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
BVH3 A66	601	KRVSFSFPPELEKK1GLINMLVKLITPDGKVLEKSGKVFGEVGVNIANFELDQPYLPQQT	660
*****			
BVH3 WU2	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
BVH3 RX1	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
BVH3 JNR7/87	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
BVH3 SP64	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
BVH3 P4241	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
BVH3 A66	661	FKYTIASKDYPPEVSYDGTFTVPTSLAYKMASQTIFYPPFHAGDTYLRVNPQFAVPKGTDAL	720
*****			
BVH3 WU2	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
BVH3 RX1	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
BVH3 JNR7/87	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
BVH3 SP64	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
BVH3 P4241	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
BVH3 A66	721	VRVFDEFHGNAYLENNYKVGEIKL1PK1LNQGTTAGNKP1PTFMANAYLDNQSTYIVE	780
*****			
BVH3 WU2	781	VPILEKENQTDKPS1LPQPKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
BVH3 RX1	781	VPILEKENQTDKPS1LPQPKRNKAQENSKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
BVH3 JNR7/87	781	VPILEKENQTDKPS1LPQPKRNKAQENKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
BVH3 SP64	781	VPILEKENQTDKPS1LPQPKRNKAQENSKLDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
BVH3 P4241	781	VPILEKENQTDKPS1LPQPKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
BVH3 A66	781	VPILEKENQTDKPS1LPQPKRNKAQENSKFDEKVEEPKTSEKVEKEKLSETGNSTSNSTL	840
*****			
BVH3 WU2	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
BVH3 RX1	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
BVH3 JNR7/87	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
BVH3 SP64	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
BVH3 P4241	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
BVH3 A66	841	EEVPTVDPVQEKVAKFAESYGMKLENVLFNMDGTIELYLPSGEVIKKNMADPTGEAPQGN	900
*****			
BVH3 WU2	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
BVH3 RX1	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
BVH3 JNR7/87	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
BVH3 SP64	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
BVH3 P4241	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
BVH3 A66	901	GENKPSENGKVSTGTVENQPTENKPADSLPEAPNEKPKPENSTDNGMLNPEGNVGSDPM	960
*****			
BVH3 WU2	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
BVH3 RX1	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
BVH3 JNR7/87	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
BVH3 SP64	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
BVH3 P4241	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
BVH3 A66	961	LDPALEEAPAVDPVQEKKLEKFTASYGLGLDSVIFNMDGTIELRLPSGEVIKKNLSDLIA	1019
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FIGURE 11

BVH11-2 SP64	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11-2 JNR7/87	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11-2 P4241	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11-2 A66	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11-2 WU2	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11-2 Rx1	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 P4241	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 WU2	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 A66	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 Rx1	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 JNR7/87	1	CSYELGRHQAGQDKKESNRVAYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 SP63	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	60
BVH11 SP64	1	CSYELGRHQAGQVKESNRVSYIDGDQAGQKAENLTPDEVSREGINAEQIVIKITDQGY	59

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BVH11-2 SP64	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK	120
BVH11-2 JNR7/87	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVDGKYYVYLK	120
BVH11-2 P4241	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11-2 A66	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11-2 WU2	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11-2 Rx1	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 P4241	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 WU2	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 A66	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 Rx1	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 JNR7/87	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 SP63	61	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	120
BVH11 SP64	60	VTSHGDHYHYYNGKPVYDAIISELLMKDPNYQLKDSDIVNEIKGGYVIKVGKYYVYLK	119

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BVH11-2 SP64	121	DAAHADNIRTKEEIKRQKQEHSHHNHSRA--DNAVAARAQGRYTTDDGYIFNASDIIE	177
BVH11-2 JNR7/87	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11-2 P4241	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11-2 A66	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11-2 WU2	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11-2 Rx1	121	DAAHADNIRTKEEIKRQKQEHSHHNHSRA--DNAVAARAQGRYTTDDGYIFNASDIIE	177
BVH11 P4241	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11 WU2	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11 A66	121	DAAHADNIRTKEEIKRQKQEHSHNHGGGSN--DQAVVAARAQGRYTTDDGYIFNASDIIE	178
BVH11 Rx1	121	DAAHADNIRTKEEIKRQKQEHSHHNHSRA--DNAVAARAQGRYTTDDGYIFNASDIIE	177
BVH11 JNR7/87	121	DAAHADNIRTKEEIKRQKQEHSHHNHSRA--DNAVAARAQGRYTTDDGYIFNASDIIE	177
BVH11 SP63	121	DAAHADNIRTKEEIKRQKQEHSHHNHSRA--DNAVAARAQGRYTTDDGYIFNASDIIE	177
BVH11 SP64	120	DAAHADNVRTKEEINRQKQEHSHQREGGTANDGAVAFARSQGRYTTDDGYIFNASDIIE	179

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BVH11-2 SP64	178	DTGDAYIVPHGDHYHYIPKNELSASELAAAAYWNGKQGSRPSSSSSYNANPVQRLSEN	237
BVH11-2 JNR7/87	179	DTGDAYIVPHGDHYHYIPKNELSASELAAAAYWNGKQGSRPSSSSSYNANPAQRLSEN	238
BVH11-2 P4241	179	DTGDAYIVPHGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11-2 A66	179	DTGDAYIVPHGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11-2 WU2	179	DTGDAYIVPRGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11-2 Rx1	178	DTGDAYIVPHGDHYHYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	237
BVH11 P4241	179	DTGDAYIVPHGDHYHYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11 WU2	179	DTGDAYIVPHGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11 A66	179	DTGDAYIVPHGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	238
BVH11 Rx1	178	DTGDAYIVPHGDHYHYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	237
BVH11 JNR7/87	178	DTGDAYIVPHGDHYHYIPKSDSLASELAAAAYWNGKQGSRPSSSSSYNANPAQRLSEN	237
BVH11 SP63	178	DTGDAYIVPHGNHFPYIPKSDSLASELAAAAYWNGKQGSRPSSSSSHNANPAQRLSEN	237
BVH11 SP64	180	DTGDAYIVPHGDHYHYIPKNELSASELAAAFLSGRNLSNLRTYRRQNSDNTPRTNWV	239

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BVH11-2 SP64	238	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	285
BVH11-2 JNR7/87	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11-2 P4241	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11-2 A66	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11-2 WU2	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11-2 Rx1	238	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	285
BVH11 P4241	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11 WU2	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11 A66	239	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	286
BVH11 Rx1	238	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	285
BVH11 JNR7/87	238	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	285
BVH11 SP63	238	HNLTVTPTYHQN-----QGENISSLLRELYAKPLSERHVESDGLIFDPAQITS	285
BVH11 SP64	240	PSVSPGTTNTNTSNNNTSQASQNSNDSSLKQLPKLPSQRHVESDGLIFDPAQITS	299

BVH11-2 SP64	286	RTARGVAVPHGNHYHFIPIYEQMSLEKRIARIIPLRYSRSHWVPSRPEQSPSQSTPEPS	345
BVH11-2 JNR7/87	287	RTARGVAVPHGNHYHFIPIYEQMSLEKRIARIIPLRYSRSHWVPSRPEQSPSQSTPEPS	346
BVH11-2 P4241	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11-2 A66	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11-2 WU2	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11-2 Rx1	286	RTANGVAVPHGDHYHFIPIYSQLSPLEEKLARIIPLRYSRSHWVPSRPEQSPSQSTPEPS	345
BVH11 P4241	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11 WU2	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11 A66	287	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQ---PS	342
BVH11 Rx1	286	RTANGVAVPHGDHYHFIPIYSQLSPLEEKLARIIPLRYSRSHWVPSRPEQSPSQSTPEPS	345
BVH11 JNR7/87	286	RTARGVAVPHGNHYHFIPIYEQMSLEKRIARIIPLRYSRSHWVPSRPEEPPSPOPTPEPS	345
BVH11 SP63	286	RTARGVAVPHGNHYHFIPIYEQMSLEERIARIIPLRYSRSHWVPSRPEQSPSQSTPEPS	345
BVH11 SP64	300	RTARGVAVPHGNHYHFIPIYEQMSLEKRIARIIPLRYSRSHWVPSRPEEPPSPOPTPEPS	359

BVH11-2 SP64	346	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	405
BVH11-2 JNR7/87	347	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	406
BVH11-2 P4241	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11-2 A66	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11-2 WU2	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11-2 Rx1	346	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	405
BVH11 P4241	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11 WU2	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11 A66	343	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	402
BVH11 Rx1	346	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	405
BVH11 JNR7/87	346	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	399
BVH11 SP63	346	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	405
BVH11 SP64	360	PSLQPAPNPQPAPSNPIDEKLVKEAVRKVGDGYVFEENGVSRYI	419

BVH11-2 SP64	406	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEVLDNLLERL	465
BVH11-2 JNR7/87	407	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	466
BVH11-2 P4241	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11-2 A66	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11-2 WU2	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11-2 Rx1	406	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	465
BVH11 P4241	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11 WU2	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11 A66	403	LAKQESLSHKLGTKCTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	462
BVH11 Rx1	406	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	465
BVH11 JNR7/87	400	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	459
BVH11 SP63	406	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	465
BVH11 SP64	420	LAKQESLSHKLGAKKTDLPSDREFYNKAYDLLARIHQDLDLNKGRQVDPEALDNLLERL	479

BVH11-2 SP64	466	KDVSSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	525
BVH11-2 JNR7/87	467	KDVPSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	526
BVH11-2 P4241	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11-2 A66	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11-2 WU2	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11-2 Rx1	466	KDVSSDKVVLVDDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	525
BVH11 P4241	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11 WU2	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11 A66	463	KDVSSDKVVLVEDILAFLAPIRHPERLGKPNQSITYTDDDEIQVAKLAGKYTTEDGYIFDP	522
BVH11 Rx1	466	KDVSSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	525
BVH11 JNR7/87	460	KDVSSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	519
BVH11 SP63	466	EDVPSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	525
BVH11 SP64	480	KDVSSDKVVLVDDILAFLAPIRHPERLGKPNQAQITYTDDDEIQVAKLAGKYTTEDGYIFDP	539

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BVH11-2 SP64	526	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	585
BVH11-2 JNR7/87	527	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	586
BVH11-2 P4241	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11-2 A66	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11-2 WU2	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11-2 Rx1	526	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	585
BVH11 P4241	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11 WU2	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11 A66	523	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	582
BVH11 Rx1	526	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	585
BVH11 JNR7/87	520	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	579
BVH11 SP63	526	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	585
BVH11 SP64	540	RDITSDEGDAYVTPHMTHSHWIKKDSLSEAERAAAQAYAKEKGLTPPSTDHQDSGNTEAK	599

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BVH11-2 SP64	586	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	645
BVH11-2 JNR7/87	587	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	646
BVH11-2 P4241	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11-2 A66	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11-2 WU2	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11-2 Rx1	586	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	645
BVH11 P4241	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11 WU2	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11 A66	583	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	642
BVH11 Rx1	586	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	645
BVH11 JNR7/87	580	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	639
BVH11 SP63	586	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	645
BVH11 SP64	600	GAEAIYNRVKAACKVPLDRMPYNLQYTVEVKNGSLIIPHVDHYHNIKFEWFDEGLYEAPK	659

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BVH11-2 SP64	646	GYSLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	690
BVH11-2 JNR7/87	647	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----VDQDSK	691
BVH11-2 P4241	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11-2 A66	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11-2 WU2	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11-2 Rx1	646	GYSLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNKNGQADTNQTEKPNEEKPQTEK	705
BVH11 P4241	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11 WU2	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11 A66	643	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----ADQDSK	687
BVH11 Rx1	646	GYSLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----NGQ	687
BVH11 JNR7/87	640	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----NGQ	681
BVH11 SP63	646	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----NGQ	687
BVH11 SP64	660	GYTLEDLLLATVKYYVEHPNERPHSDNGPGN ASDHVRKNK-----NGQ	701

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BVH11-2 SP64	691	PDEDKEHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	750
BVH11-2 JNR7/87	692	PDEDKEHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	751
BVH11-2 P4241	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11-2 A66	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11-2 WU2	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11-2 Rx1	706	PEEDKEHDEVSEPTHPESDEKENHAGVLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11 P4241	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	765
BVH11 WU2	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11 A66	688	PDEDKGHDEVSEPTHPESDEKENHAGLNPSADNLYKPSTDTEETEEEADTTDEAEIPQV	747
BVH11 Rx1	688	ADTNQTEKPNEEKPQTEKPREEETPREEKPQSEKPKPTEEEESPEESEEPQV	747
BVH11 JNR7/87	682	ADTNQTEKPNEEKPQTEKPREEETPREEKPQSEKPKPTEEEESPEESEEPQV	741
BVH11 SP63	688	ADTNQTEKPSEEKPQTEKPREEETPREEKPQSEKPKPESP---KPTEEEESPEESEEPQV	743
BVH11 SP64	702	ADTNQTEKPSEEKPQTEKPREEETPREEKPQSEKPKPESP---KPTEEEESPEESEEPQV	757
* * * * *			
BVH11-2 SP64	751	ENSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	810
BVH11-2 JNR7/87	752	ENSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	811
BVH11-2 P4241	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11-2 A66	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11-2 WU2	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11-2 Rx1	766	EYSVINAKIABAEALLEKVTDSSIRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	825
BVH11 P4241	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11 WU2	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11 A66	748	EHSVINAKIADAEEALLEKVTDPISRQNAMEETLTGKSSLLLGTKDNNNTISAEVDSLLALL	807
BVH11 Rx1	748	ETEKVEEKLREAEDLILGKIQNPPIIKSNAKETLTGLKNNNLLFGTQDNNTIMAEAEKLLALL	807
BVH11 JNR7/87	742	ETEKVEEKLREAEDLILGKIQNPPIIKSNAKETLTGLKNNNLLFGTQDNNTIMAEAEKLLALL	801
BVH11 SP63	744	ETEKVEEKLREAEDLILGKIQDPPIIKSNAKETLTGLKNNNLLFGTQDNNTIMAEAEKLLALL	803
BVH11 SP64	758	ETEKVEEKLREAEDLILGKIQDPPIIKSNAKETLTGLKNNNLLFGTQDNNTIMAEAEKLLALL	817
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BVH11-2 SP64	811	KESQPAPIQ	819
BVH11-2 JNR7/87	812	KESQPAPIQ	820
BVH11-2 P4241	808	KKSQPAPIQ	816
BVH11-2 A66	808	KKSQPAPIQ	816
BVH11-2 WU2	808	KKSQPAPIQ	816
BVH11-2 Rx1	826	KESQPAPIQ	834
BVH11 P4241	808	KESK	811
BVH11 WU2	808	KESK	811
BVH11 A66	808	KESK	811
BVH11 Rx1	808	KESK	811
BVH11 JNR7/87	802	KESK	805
BVH11 SP63	804	KESK	807
BVH11 SP64	818	KESK	821

FIGURE 12

BVH11-2	BVH11	BVH11	BVH11-2	BVH11	BVH11-2	BVH11	BVH11-2	BVH11	BVH11-2
SP63	JNR 7/87	JNR 7/87	WU2	WU2	A66	P4241	P4241	Rx-1	Rx-1
I 88%	I 88%	I 82%	I 80%	I 80%	I 80%	I 80%	I 80%	I 88%	I 81%
S 90%	S 91%	S 87%	S 85%	S 85%	S 85%	S 85%	S 85%	S 91%	S 85%
I 87%	I 87%	I 98%	I 95%	I 96%	I 95%	I 96%	I 95%	I 87%	I 94%
S 90%	S 90%	S 98%	S 96%	S 97%	S 96%	S 97%	S 97%	S 90%	S 95%
I 96%	I 96%	I 88%	I 88%	I 87%	I 88%	I 87%	I 88%	I 87%	I 89%
S 96%	S 96%	S 91%	S 91%	S 90%	S 91%	S 90%	S 91%	S 97%	S 91%
I 87%	I 87%	I 87%	I 86%	I 87%	I 86%	I 87%	I 87%	I 97%	I 89%
S 90%	S 91%	S 91%	S 90%	S 91%	S 90%	S 91%	S 90%	S 96%	S 90%
I 96%	I 96%	I 97%	I 96%	I 97%	I 96%	I 96%	I 97%	I 87%	I 94%
S 97%	S 97%	S 98%	S 97%	S 98%	S 97%	S 97%	S 98%	S 90%	S 95%
I 98%	I 98%	I 92%	I 92%	I 98%	I 99%	I 98%	I 98%	I 87%	I 94%
S 98%	S 98%	S 94%	S 94%	S 98%	S 99%	S 98%	S 98%	S 90%	S 95%
I 98%	I 98%	I 99%	I 99%	I 98%	I 99%	I 99%	I 99%	I 86%	I 93%
S 98%	S 98%	S 99%	S 99%	S 98%	S 99%	S 99%	S 99%	S 90%	S 95%
I 99%	I 99%	I 99%	I 99%	I 100%	I 99%	I 99%	I 97%	I 92%	I 92%
S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 91%	S 91%	A66	A66
I 99%	I 99%	I 99%	I 99%	I 99%	I 99%	I 86%	I 86%	I 93%	I 93%
S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 90%	S 90%	A66	A66
I 99%	I 99%	I 99%	I 99%	I 99%	I 99%	I 99%	I 99%	I 92%	I 92%
S 99%	S 99%	S 99%	S 99%	S 99%	S 99%	S 91%	S 91%	P4241	P4241
I 86%	I 86%	I 87%	I 87%	I 87%	I 87%	I 87%	I 87%	I 93%	I 93%
S 90%	S 90%	S 91%	S 91%	S 91%	S 91%	S 91%	S 91%	P4241	P4241
I 91%	I 91%	I 92%	I 92%	I 92%	I 92%	I 92%	I 92%	BVH11	BVH11
S 92%	S 92%	S 92%	S 92%	S 92%	S 92%	S 92%	S 92%	Rx-1	Rx-1

FIGURE 13

AATTCCCTTGT	CGGGTAAGTT	CCGACCCGCA	CGAAAGGCGT	AATGATTTGG	GCACTGTCTC	60
AAAGGAGAGAC	TCGGTGAAT	TTAGTACCT	GTGAAGATGC	AGTTACCCG	CGACAGGACG	120
GAAAGACCCC	ATGGAGCTTT	ACTGCAGTT	GATATTGAGT	GTCTGTACCA	CATGTACAGG	180
ATAGGTAGGA	GTCTAAGAGA	TCGGGACGCC	AGTTTCAAG	GAGACGCTGT	TGGGATACTA	240
CCCTTGTGTT	ATGGCCACTC	TAACCCAGAT	AGGTGATCCC	TATCGGAGAC	AGTGTCTGAC	300
GGGCAGTTG	ACTGGGGCGG	TCGCCCTCTA	AAAGGTAACG	GAGGCGCCCA	AAGGTTCCCT	360
CAGAATGGTT	GGAAATCATT	CCGAGAGTGT	AAAGGTATAA	GGGAGCTTGA	CTGCGAGAGC	420
TACAACATCGA	GCAGGGACGA	AAGTCGGGCT	TAGTGATCCG	GTGGTTCCGT	ATGGAAGGGC	480
CATCGCTCAA	CGGATAAAAAG	CTACCCCTGGG	GATAACAGGC	TTATCTCCCC	CAAGAGTTCA	540
CATCGACGGG	GAGGTTTGGC	ACCTCGATGT	CGGCTCGTCG	CATCCCTGGGG	CTGTAGTCGG	600
TCCCAAGGGT	TGGGCTGTT	GCCCATTAAA	CGGGCACCGCG	AGCTGGGTTTC	AGAACGTCGT	660
GAGACAGTTG	GGTCCCTATC	CGTCGCGGGC	GTAGGAAATT	TGAGAGGATC	TGCTCCTAGT	720
ACGAGAGGAC	CAGAGTGGAC	TTACCGCTGG	TGTACCAAGT	GTCTTGCCAA	AGGCATCGCT	780
GGTAGCTAT	GTAGGGAGG	GATAAACGCT	AAAGGATCT	AAGTGTGAAA	CCCACCTCAA	840
GATGAGATTT	CCCATGATTA	TATATCAGTA	AGAGCCCTGA	GAGATGATCA	GGTAGATAGG	900
TTAGAAGTGG	AAAGTGGCG	ACACATGTAG	CGGACTAATA	CTAATAGCTC	GAGGACTTAT	960
CCAAAGTAAC	TGAGAATATG	AAAGCGAACG	GTTTCTTAA	ATTGAATAGA	TATTCAATTT	1020
TGAGTAGGTA	TTACTCAGAG	TTAAGTGAAG	ATAGCCTAGG	AGATACACCT	GTACCCATGC	1080
CGAACACAGA	AGTTAACGCCC	TAGAACGCCG	GAAGTAGTTG	GGGTTGCC	CCTGTGAGAT	1140
AGGGAAGTCG	CTTAGCTCTA	GGGAGTTAG	CTCAGCTGGG	AGAGCATCTG	CCTTACAAGC	1200
AGAGGGTCAG	CGGTTCCATC	CCGTTAACTC	CCAAAGGTC	CGTAGTGTAG	CGGTTATCAC	1260
GTCGCCCTGT	CACGGCGAAG	ATCGGGGTT	CGATTCCCGT	CGGGACCGTT	TAAGGTAACG	1320
CAAGTTATTT	TAGACTCGTT	AGCTCAGTTG	GTAGAGCAAT	TGACTTTAA	TCAATGGTC	1380
ACTGGTTCGA	GCCCAGTACG	GGTCATATAT	CGGGTTTGG	CGGAATTCTA	ATCTCTTGA	1440
AATCATCTTC	TCTCACTTTC	CAAAACTCTA	TTACCTCTTA	TTATACCACA	TTTCAATCTT	1500
CAACTTCCC	GTAATATAAG	CACCTCTGGC	GAAAGAAGTT	TCAATGTCT	AAAGTAATAA	1560
GTGAATCCAA	TTCAGGAAC	CCAAGAACAA	AAGAAACATC	TGGTGTACACA	AGTATTGGAT	1620
GGCACAGAGT	CACGTGGTAG	TCTGACCCCTA	GCAGAAATT	TAAATAGTAA	ACTATTTACT	1680
GGTTAATTAA	ATGGTTAAAT	AACCGTTTA	GAAAACATT	TAATAAAGTA	AAAGAAGTTG	1740
AGAAAAAAACT	TCATCATTTA	TTGAAATGAG	GGATTATGAG	AATTAGTAA	AAAATATATA	1800
GCAGCTGGAT	CAGCTGTTAT	CGTATCCTTG	AGTCTATGTG	CCTATGCACT	AAACCAGCAT	1860
CGITCGCAGG	AAAATAAGGA	CAATAATCGT	GTCTCTTATG	TGGATGGCAG	CCAGTCAGT	1920
CAGAAAAGTG	AAAACCTGAC	ACCAGACCAG	GTAGGCCAGA	AAGAAGGAAT	TCAGGCTGAG	1980
CAAATTGTA	TCAAAATTAC	AGATCAGGGC	TATGTAACGT	CACACGGTGA	CCACTATCAT	2040
TACTATAATG	GGAAAGTTCC	TTATGATGCC	CTCTTTAGTG	AAGAACTCTT	GATGAAGGAT	2100
CCAAACTATC	AACTTAAAGA	CGCTGATATT	GTCAATGAAG	TCAAGGGTGG	TTATATCATC	2160
AAGGTCGATG	GAAAATATTA	TGTCTACCTG	AAAGATGCAG	CTCATGCTGA	TAATGTTCGA	2220
ACTAAAGATG	AAATCAATCG	TCAAAAACAA	GAACATGTCA	AAGATAATGA	GAAGGTTAAC	2280
TCTAATGTTG	CTGTAGCAAG	GTCTCAGGGG	CGATATACGA	CAAATGATGG	TTATGTCITT	2340
AATCCAGCTG	ATATTATCGA	AGATACGGGT	AATGTTATA	TCGTTCTCA	TGGAGGTCAC	2400
TATCACTACA	TTCCCAAAG	CGATTATCT	GCTAGTGAAT	TAGCAGCAGC	TAAAGCACAT	2460
CTGGCTGGAA	AAAATATGCA	ACCGAGTCAG	TTAAGCTATT	CTTCAACAGC	TAGTGACAAT	2520
AACACGCAAT	CTGTAGAAA	AGGATCAACT	AGCAAGCCAG	CAAATAATC	TGAAAATCTC	2580
CAGAGTCTTT	TGAAGGAAC	CTATGATCA	CCTAGCGCCC	AACGTTACAG	TGAATCAGAT	2640
GGCCTGGTCT	TTGACCCCTGC	TAAGATTATC	AGTCGTACAC	CAAATGGAGT	TGCGATTCCG	2700
CATGGCGACC	ATTACCACT	TATTCCTTAC	AGCAAGCTTT	CTGTTTACA	AGAAAAGATT	2760
GCCAGAATGG	TGCCCTATCAG	TGGAATGGT	TCTACAGTTT	CTACAAATGC	AAAACCTAA	2820
GAAGTAGTGT	CTAGTCTAGG	CAGCTTTCA	AGCAATCTT	CTTCTTAAAC	GACAAGTAAG	2880
GAGCTCTT	CAGCATCTGA	TGGTTATATT	TTAATCCAA	AAGATATCGT	TGAAGGAAACG	2940
GCTACAGOTT	ATATTGTAAG	ACATGGTGAT	CATTCTCATT	ACATTCCAAA	ATCAAATCAA	3000
ATTGGGCAAC	CGACTCTTCC	AAACAATAGT	CTAGCAACAC	CTTCTCCATC	TCTTCCAATC	3060
AATCCAGGAA	CTTCACATGA	AAAACATGAA	GAAGATGGAT	ACGGATTGTA	TGCTAATCGT	3120
ATTATCGCTG	AAGATGAATC	AGGTTTGTG	ATGAGTCACG	GAGACACAA	TCATTATTT	3180
TTCAAGAAGG	ACTTGACAGA	AGAGCAAATT	AAAGGCTGCGC	AAAACATT	AGAGGAAGTT	3240
AAAACATGTC	ATAATGGATT	AGATTCTTG	TCATCTCATG	AACAGGATTA	TCCAGGTAAT	3300
GCCAAAGAAA	TGAAAGATTT	AGATAAAAAA	ATCGAAGAAA	AAAATGCTGG	CATTATGAAA	3360

CAATATGGTG	TCAAACGTGA	AA GTATTGTC	GTGAATAAG	AAAAAAATGC	GATTATTTAT	3420
CCGCATGGAG	ATCACCACCA	TGCAGATCCG	ATTGATGAAC	ATAAACCGGT	TGGAATTGGT	3480
CATTCTCACA	GTAACTATGA	ACTGTTAAA	CCCGAAGAAG	GAGTTGCTAA	AAAAGAAGGG	3540
AATAAAGTT	ATACTGGAGA	AGAATTAACG	AATGTTGTTA	ATTGTTAAA	AAATAGTACG	3600
TTTAATAATC	AAAACCTTAC	TCTAGCCAAT	GGTCAAAAAC	GCGTTTCTTT	TAGTTTCCG	3660
CCTGAATTGG	AGAAAAAAATT	AGGTATCAAT	ATGCTAGTAA	AATAAATAAC	ACCAGATGGA	3720
AAAGTATTGG	AGAAAGTATC	TGGTAAAGTA	TTTGGAGAAG	GAGTAGGGAA	TATTGCAAAC	3780
TTTGAATTAG	ATCAACCTTA	TTTACCCAGGA	CAAACATTTA	AGTATACTAT	CGCTTCAAAAA	3840
GATTATCCAG	AA GTAAGTTA	TGATGGTACA	TTTACAGTTC	CAACCTCTT	AGCTTACAAA	3900
ATGGCCAGTC	AAACGATTTT	CTATCCTTTC	CATGCAGGGG	ATACTTATT	AAGAGTGAAC	3960
CCTCAATTG	CA GTGCCTAA	AGGAACTGAT	GCTTAGTCA	GAGTGTTGA	TGAATTTCAT	4020
GAAAATGTT	ATTTAGAAAA	TAACTATAAA	GTGGTAAA	TCAAAATTAC	GATTCCGAAA	4080
TTAAACCAAG	GAACAACCAAG	AACGGCCGGA	AATAAAATTC	CTGTAACCTT	CATGGCAAAT	4140
GCTTATTG	ACAATCAATC	GACTTATATT	GTGGAAGTAC	CTATCTTGG	AAAAGAAAAAT	4200
CAAACGTATA	AA CCAAGTAT	TCTACCACAA	TTTAAAAGGA	ATAAAGCACA	AGAAAAACTCA	4260
AAACTTGATG	AAAAGGTAGA	AGAACCAAAG	ACTAGTGAGA	AGGTAGAAA	AGAAAAACTT	4320
TCTGAAACTG	GGAAATAGTAC	TAGTAATTCA	ACGTTAGAAG	AGTTCCCTAC	AGTGGATCCT	4380
GTACAAGAAA	AA GTAGAAA	ATTGCTGAA	AGTTATGGG	TGAAGCTAGA	AAATGTCTTG	4440
TTTAATATGG	ACGGAACAAT	TGAATTATAT	TTACCATCAG	GAGAAGTCAT	AAAAAAGAAT	4500
ATGGCAGATT	TTACAGGAGA	ACCACCTCA	GGAAATGGT	AAAATAAAC	ATCTGAAAAT	4560
GGAAAAGTAT	CTACTGGAAC	AGTTGAGAAC	CAACCAACAG	AAAATAAAC	AGCAGATTCT	4620
TTACCCAGAGG	CACCAAACGA	AAAACCTGTA	AAACCAGAAA	ACTAACCGGA	TAATGGAATG	4680
TTGAATCCAG	AA GGGATGT	GGGGAGTGAC	CCTATGTTAG	ATCCAGCATT	AGAGGAAGCT	4740
CCACGAGTAG	ATCCGTACA	AGAAAAATT	AAAAATT	CAGCTAGTTA	CGGATTAGGC	4800
TTAGATAGTG	TTATATTCAA	TATGGATGGA	ACGATTGAAT	TAAGATTGCC	AA GTGGAGAA	4860
GTGATAAAA	AGAATTATC	TGATTTCTATA	GCGTAAGGAA	TAGCAGTAGA	AAAAGTCTGA	4920
ATCAAAAATG	AA GTTCTCTC	AAAGTTAGA	AATAAAACTC	TGACTTTGGG	AGAATTTCAT	4980
TTTATTATTA	ATATATAAA	TTTCTTGACA	TACAACTTAA	AAAGAGGTGG	AATATTACT	5040
AGTTAATT	(SEQ ID NO : 11)					5048

FIGURE 14

CAGAGATCTT	AGTGAATCAA	ATATACTTAA	AAAAAGAGGA	AAGAATGAAA	ATCAATAAAA	60
AATATCTAGC	TGGGTCACTA	GCTACACTTG	TTTTAAGTGT	CTGTGCTTAT	GAACTAGGTT	120
TGCATCAAGC	TCAAACGTAA	AAAGAAAATA	ATCGTGTTC	CTATATAGAT	GGAAAACAAG	180
CGACGCAAA	AACGGAGAAT	TTGACTCTG	ATGAGGTAG	CAAGCGTGA	GGAAATCAACG	240
CCGAACAAAT	CGTCATCAAG	ATTACGGATC	AAGGTTATGT	GACCTCTCAT	GGAGACCATT	300
ATCATTACTA	TAATGGCAAG	GTCCCTTATG	ATGCCATCAT	CAGTGAAGAG	CTCCTCATGA	360
AAGATCCGAA	TTATCAGTTG	AAGGATTCA	ACATTGTCAA	TGAAATCAAG	GGTGGTTATG	420
TCATTAAGGT	AAACGGTAAA	TACTATGTT	ACCTTAAGGA	TGCAAGCTCAT	GCGGATAATG	480
TCCGTACAAA	AGAAGAAATC	AATCGGAAA	AACAAGAAC	TAGTCAGCAT	CGTGAAGGAG	540
GGACTTCAGC	AAACGATGGT	GCGGTAGCT	TTGCACGTC	ACAGGGACGC	TACACCACAG	600
ATGATGGTTA	TATCTCAAT	GCATCTGATA	TCATCGAAGA	TACGGGCGAT	GCCTATATCG	660
TTCCCTCATGG	AGATCATTAC	CATTACATC	CTAAGAACATG	GTTATCAGCT	AGCGAGTTGG	720
CTGCTGCAGA	AGCCTCCCTA	TCTGGTCGGG	AAAATCTGTC	AAATTAAAGA	ACCTATGCC	780
GACAAAATAG	CGATAACACT	CCAAGAACAA	ACTGGGTACC	TTCTGTAAAG	AATCCAGGAA	840
CTACAAATAC	TAACACAAGC	AACAACAGCA	ACACTAACAG	TCAAGCAAGT	CAAAGTAATG	900
ACATTGATAG	TCTCTGAAA	CAGCTCTACA	AACTGCCCTT	GAGTCAACGC	CATGTAGAAT	960
CTGATGGCCT	TATTTTCGAC	CCAGCGAAA	TCACAAGTCG	AACCGCCAGA	GGTGTAGCTG	1020
TCCCTCATGG	TAACCATTAC	CACTTTATCC	CTTATGAACA	AATGTCGAA	TTGGAAAAAC	1080
GAATTGCTCG	TATTAATCCC	CTTCGTTATC	GTTCAAACCA	TTGGGTACCA	GATTCAAGAC	1140
CAGAAGAACCC	AAAGTCCACAA	CCGACTCCAG	AACTTAGTCC	AAAGTCCGCAA	CCTGCACCAA	1200
ATCCTCAACC	AGCTCCAAGC	AATCCAATTG	ATGAGAAATT	GGTCAAAGAA	GCTGTTGAA	1260
AAGTAGGCAGA	TGGTTATGTC	TTTGAGGAGA	ATGGAGTTTC	TCGTTATATC	CCAGCCAAGA	1320
ATCTTCAGC	AGAAACAGCA	GCAGGCATTG	ATAGCAAAC	GGCCAAGCAG	GAAAGTTAT	1380
CTCATAAGCT	AGGAGCTAAG	AAAACGTGACC	TCCCCATCTAG	TGATCGAGAA	TTTTACAATA	1440
AGGCTTATGA	CTTACTAGCA	AGAATTCA	AAGATTTACT	TGATAATAAA	GGTCGACAAG	1500
TTGATTTTGAT	GGCTTTGGAT	AACCTGTTGG	AACGACTCAA	GGATGTCCTCA	AGTGTAAAG	1560
TCAAGTTAGT	GGATGATATT	CTTGCCTTCT	TAGCTCCGAT	TCGTCATCCA	GAACGTTTAG	1620
GAAAACCAAA	TGCGCAAATT	ACCTACACTG	ATGATGAGAT	TCAAGTAGCC	AAGTTGGCAG	1680
GCAAGTACAC	AACAGAAGAC	GGTTATATCT	TTGATCCTCG	TGATATAACC	AGTGTAGAGG	1740
GGGATGCTA	TGTAACTCCA	CATATGACCC	ATAGCCACTG	GATTAAAAAA	GATAGTTTGT	1800
CTGAGCTGA	GAGAGCGGCA	GCCCAGGCTT	ATGCTAAAGA	GAAAGGTTTG	ACCCCTCCTT	1860
CGACAGACCA	TCAGGATTCA	GGAAATACTG	AGGCAAAAGG	AGCAGAACG	ATCTACAACC	1920
GCGTGAAGC	AGCTAAGAAG	GTCGCACTTG	ATCGTATGCC	TTACAATCTT	CAATATACTG	1980
TAGAAGTCAA	AAACGGTAGT	TTAACATAC	CTCATTATGA	CCATTACCAT	ACATCAAAT	2040
TTGAGTGGTT	TGACGAAGGC	CTTTATGAGG	CACCTAAGGG	GTATACTCTT	GAGGATCTTT	2100
TGGCAGCTGT	CAAGTACTAT	GTCGAACATC	CAAACGAACG	TCCGCATTCA	GATAATGGTT	2160
TTGGTAACGC	TAGCGACCAT	GTTCAAAGAA	ACAAAAATGG	TCAAGCTGAT	ACCAATCAA	2220
CGGAAAAACC	AAGCGAGGAG	AAACCTCAGA	CAGAAAAACCC	TGAGGAAGAA	ACCCCTCGAG	2280
AAGAGAAACC	ACAAACCGAG	AAACCAGAGT	CTCCAAAACC	AACAGAGGAA	CCAGAAGAAG	2340
AATCACCAGA	GGAAATCAGAA	GAACCTCAGG	TCGAGACTGA	AAAGGTTGAA	AAAAAACTGA	2400
GAGAGGCTGA	AGATTTACTT	GGAAAAATCC	AGGATCAAAT	TATCAAGTCC	AATGCCAAAG	2460
AGACTCTCAC	AGGATTTAAA	AATAATTAC	TATTTGGCAC	CCAGGACAAC	AATACTATTA	2520
TGGCAGAACG	TGAAAAAAACTA	TTGGCTTTAT	TAAGGAGAG	TAAGTAAAGG	TAGCAGCATT	2580
TTCTAATCC	TAACAAACAGG	ATAGGAGAAC	GGGAAAACGA	AAAATGAGAG	CAGAATGTGA	2640
TTTCTAG	(SED ID NO : 12)					2647

FIGURE 15

GGGTCTTAAA	ACTCTGAATC	CTTGTAGAGGC	AGACCCACAA	AATGACAAGA	CCTATTTAGA	60
AAATCTGGAA	AAAAATATGA	GTGTTCTAGC	AGAAGAATTA	AACTGAGGAA	AGAATGAAAA	120
TCAATAAAA	ATATCTAGCA	GTTCAGTGG	CAGTCCTTGC	CCTAAGTGTGTT	TGTTCTATG	180
AACTTGGTCG	TCACCAAGCT	GTCAGGTAA	AGAAAGAGTC	TAATCGAGTT	TCTTATATAG	240
ATGGTGTATCA	GGCTGGTCAA	AAGGCAGAAA	ATTTGACACC	AGATGAAGTC	AGTAAGAGAG	300
AGGGGATCAA	CGCCGAACAA	ATTGTTATCA	AGATTACGGA	TCAAGGTTAT	GTGACCTCTC	360
ATGGAGACCA	TTATCATTAC	TATAATGGCA	AGGTTCCCTA	TGATGCCATC	ATCAGTGAAG	420
AACTTCTCAT	GAAAGATCCG	ATTATTCAGT	TGAAGGATTC	AGACATTGTC	AATGAAATCA	480
AGGGTGGCTA	TGTGATTAAG	GTAGACGGAA	AATACTATGT	TTACCTTAAA	GATGCGGCC	540
ATGCGGACAA	TATTGGACAA	AAAGAAGAGA	TTAAACGTCA	GAAGCAGGAA	CACAGTCATA	600
ATCATAACTC	AAGAGCAGAT	AATGCTGTTG	CTGCAGCCAG	AGCCCAAGGA	CGTTATACAA	660
CGGATGATGG	GTATATCTTC	AATGCATCTG	ATATCATGTA	GGACACGGGT	GATGCTTATA	720
TCGTTCTCA	CGGCGACCAT	TACCATTACA	TTCTTAAGAA	TGAGTTATCA	GCTAGCGAGT	780
TAGCTGCTGC	AGAACGCTAT	TGGAATGGGA	AGCAGGGATC	TCGTCCTTCT	TCAAGTTCTA	840
GTTATAATGC	AAATCCAGTT	CAACCAAGAT	TGTCAGAGAA	CCACAATCTG	ACTGTCACTC	900
CAACTTATCA	TCAAAATCAA	GGGGAAAACA	TTTCAAGCCT	TTTACGTGAA	TTGTATGCTA	960
AACCCCTTATC	AGAACGCCAT	GTAGAACTG	ATGGCCTTAT	TTTCGACCCA	GCGAAATCA	1020
CAAGTCGAAC	CGCCAGAGGT	GTAGCTGTCC	CTCATGGTAA	CCATTACAC	TTTATCCCTT	1080
ATGAACAAAT	GTCTGAATTG	GAAAAACGAA	TTGCTCGTAT	TATTCCTCTT	CGTTATCGTT	1140
CAAACCATTG	GGTACCAAGAT	TCAAGACCAAG	AACAACCAAG	TCCACAATCG	ACTCCGGAAC	1200
CTAGTCCAAG	TCTGCAACCT	GCACCAAATC	CTCAACCCAGC	TCCAAGCAAT	CCAATTGATG	1260
AGAAAATTGGT	CAAAGAAGCT	GTTGAAAAG	TAGGCGATGG	TTATGTCTTT	GAGGAGAATG	1320
GAGTTTCTCG	TTATATCCCA	GCCAAGGATC	TTTCAGCAGA	AACAGCAGCA	GGCATTGATA	1380
GCAAACTGGC	CAAGCAGGAA	AGTTTATCTC	ATAAGCTAGG	AGCTAAGAAA	ACTGACCTCC	1440
CATCTAGTGA	TCGAGAATT	TACAATAAGG	CTTATGACTT	ACTAGCAAGA	ATTCAACCAAG	1500
ATTTACTTGA	TAATAAAAGGT	CGACAAGTTG	ATTTTGAGGT	TTTGATAAC	CTGTTGGAAC	1560
GACTCAAGGA	TGTCTCAAGT	GATAAAAGTC	AGTTAGTGG	TGATATTCTT	GCCTTCCTAG	1620
CTCCGATTGCG	TCATCCAGAA	CGTTTAGGAA	AACCAAATGC	GCAAAATTAC	TACACTGATG	1680
ATGAGATTCA	AGTAGCCAAG	TTGGCAGGCA	AGTACACAAAC	AGAACAGGT	TATATCTTTG	1740
ATCCTCGTGA	TATAACCAGT	GATGAGGGGG	ATGCCTATGT	AACTCCACAT	ATGACCCATA	1800
GCCACTGGAT	AAAAAAAGAT	AGTTTGTCTG	AAGCTGAGAG	AGCAGCAGCC	CAGGCTTATG	1860
CTAAAGAGAA	AGGTTTGACC	CCTCTTCGA	CAGACCACCA	GGATTCAAGGA	AATACTGAGG	1920
CAAAAGGAGC	AGAAGCTATC	TACAACCGCG	TGAAAGCAGC	TAAGAAGGTG	CCACTTGATC	1980
GTATGCCCTA	CAATCTTCAA	TATACTGTAG	AAGTCAAAAAA	CGGTAGTTTA	ATCATACCTC	2040
ATTATGACCA	TTACCATAAC	ATCAAATTG	AGTGGTTTGA	CGAAGGCCCTT	TATGAGGCAC	2100
CTAAGGGTA	TAGTCTTGAG	GATCTTTGG	CGACTGTCAA	GTACTATGTC	GAACATCCAA	2160
ACGAACGTCC	GCATTCAGAT	AATGGTTTG	GTAACGCTAG	TGACCATGTT	CGTAAAAATA	2220
AGGCAGACCA	AGATAGTAA	CCTGATGAAG	ATAAGGAACA	TGATGAAGTA	AGTGAGCCAA	2280
CTCACCCCTGA	ATCTGATGAA	AAAGAGAAC	ACGCTGGTTT	AAATCCTTCA	GCAGATAATC	2340
TTTATAAACC	AAGCACTGAT	ACGGAAGAGA	CAGAGGAAGA	AGCTGAAGAT	ACCACAGATG	2400
AGGCTGAAAT	TCCTCAAGTA	GAGAATTCTG	TTATTAACGC	TAAGATAGCA	GATGCGGAGG	2460
CCTTGCTAGA	AAAAGTAACA	GATCCTAGTA	TTAGACAAA	TGCTATGGAG	ACATTGACTG	2520
GTCTAAAAG	TAGTCTTCTT	CTCGAACGA	AAGATAATAA	CACTATTCA	GCAGAAGTAG	2580
ATAGTCTCTT	GGCTTTGTTA	AAAGAAAAGTC	AACCGGCTCC	TATACAGTAG	TTAAATGAA	2639

(SEQ ID NO : 13)

FIGURE 16

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MKINKKYL	AG SVAVLALSVC SYELGRHQAG QVKKESNRVS YIDGDQAGQK	50
AENLTPDEVS	KREGINAEQI VIKITDQGYV TSHGDHYHYY NGKVPYDAII	100
SEELLMKDPN	YQLKDSDIVN EIKGGYVIKV DGKYYVYLKD AAHADNIRTK	150
EEIKRQKQEH	SHNHNSRADN AVAAARAQGR YTTDDGYIFN ASDIIIEDTGD	200
AYIVPHGDHY	HYIPKNELSA SELAAAEEAYW NGKQGSRPSS SSSYNANPVQ	250
PRLSENHNL	VTPTYHQNQG ENISSLLREL YAKPLSERHV ESDGLIFDPA	300
QITSRTARGV	AVPHGNHYHF IPYEQMSELE KRIARIIPLR YRSNHWVPDS	350
RPEQPSPOST	PEPSPSLQPA PNPNQAPSNP IDEKLVKEAV RKVGDGYVFE	400
ENGVSRYIPA	KDLSAETAAG IDSKLAKQES LSHKLGAKKT DLPSSTDREFY	450
NKAYDLLARI	HQDLLDNKGR QVDFEVLDNL LERLKDVSSD KVKLVDDILA	500
FLAPIRHPER	LGKPNAQITY TDDEIQVAKL AGKYTTEDGY IFDPRDITSD	550
EGDAYVTPHM	THSHWIKKDS LSEAERAQAAQ AYAKEGLTP PSTDHQDSGN	600
TEAKGAEAIY	NRVKAACKVP LDRMPYNLQY TVEVKNGSLI IPHYDHYHNI	650
KFEWFDEGLY	EAPKGYSLED LLATVKYYVE HPNERPHSDN GFGNASDHVR	700
KNKADQDSKP	DEDKEHDEVS EPTHPESDEK ENHAGLNPSA DNLYKPSTDT	750
EETEEEAEADT	TDEAEIPQVE NSVINAKIAD AEALLEKVTD PSIRQNAME	800
LTGLKSSLLL	GTKDNNTISA EVDSLALLK ESQPAPIQ	838
(SEQ ID NO : 14)		

FIGURE 17

TGTGCCTATG	CACTAAACCA	GCATCGTTCG	CAGGAAAATA	AGGACAATAA	TCGTGTCTCT	60
TATGTGGATG	GCAGCCAGTC	AAAGTCAGAAA	AGTGAAAACT	TGACACCAGA	CCAGGTTAGC	120
CAGAAAAGAAG	GAATTCAAGG	TGAGCAAATT	GTAATCAAAA	TTACAGATCA	GGGCTATGTA	180
ACGTACACAG	GTGATCACTA	TCATTACTAT	AATGGGAAAG	TTCCCTATGA	TGCCCTCTTT	240
AGTGAAGAAC	TCTTGATGAA	GGATCCAAAC	TATCAACTTA	AAGACGCTGA	TATTGTCAT	300
GAAGTCAAGG	GTGGTTATAT	CATCAAGGTC	GATGGAAAAT	ATTATGTCTA	CCTGAAAGAT	360
GCAGCTCATG	CTGATAATGT	TCGAACAAA	GATGAAATCA	ATCGTCAAAA	ACAAGAACAT	420
GTCAAAGATA	ATGAGAAGGT	TAACTCTAAT	GTGCTGTAG	CAAGGTCTCA	GGGACGATAT	480
ACGACAAATG	ATGGTTATGT	CTTAAATCCA	GCTGATATTA	TCGAAGATAC	GGGTAATGCT	540
TATATCGTTC	CTCATGGAGG	TCACTATCAC	TACATTCCA	AAAGCGATT	ATCTGCTAGT	600
GAATTAGCAG	CAGCTAAAGC	ACATCTGGCT	GGAAAAAATA	TGCAACCGAG	TCAGTTAACG	660
TATTCTCAA	CACCTCTCC	ATCTCTTCCA	ATCAATCCAG	GAACCTCACA	TGAGAAACAT	720
GAAGAAGATG	GATACGGATT	TGATGCTAAT	CGTATTATCG	CTGAAGATGA	ATCAGGTTTT	780
GTCATGAGTC	ACGGAGACCA	CAATCATTAT	TTCTTCAAGA	AGGACTTGAC	AGAAGAGCAA	840
ATTAAGGCTG	CGCAAAAACA	TTTAGAGGAA	GTAAAACTA	GTCTATAATGG	ATTAGATTCT	900
TTGTCATCTC	ATGAAACAGGA	TTATCCAAGT	AATGCCAAAG	AAATGAAAGA	TTTAGATAAA	960
AAAATCGAAG	AAAAAATTCG	TGGCATTATG	AAACAATATG	GTGCTAAACG	TGAAAGTATT	1020
GTCGTGAATA	AAGAAAAAAA	TGGCATTATT	TATCCGATG	GAGATCACCA	TCATGCAGAT	1080
CCGATTGATG	AACATAAAACC	GGTGGAAATT	GGTCATTCTC	ACAGTAACTA	TGAACGTGTT	1140
AAACCCGAAG	AAGGAGTTGC	TTAAAAAGAA	GGGAATAAAG	TTTATACTGG	AGAAGAATTA	1200
ACGAATGTTG	TTAATTGTT	AAAAAATAGT	ACGTTTAATA	ATCAAAACTT	TACTCTAGCC	1260
AATGGTCAAA	AACCGCTTTC	TTTAGTTTT	CCGCCTGAAT	TGGAGAAAAA	ATTAGGTATC	1320
AATATGCTAG	TTAAATTAAAT	AACACCAGAT	GGAAAAGTAT	TGGAGAAAGT	ATCTGGTAAA	1380
GTATTTGGAG	AAGGAGTAGG	GAATATTGCA	AACTTTGAAT	TAGATCAACC	TTATTTACCA	1440
GGACAAACAT	TTAAGTATAC	TATCGCTTCA	AAAGATTATC	CAGAAGTAAG	TTATGATGGT	1500
ACATTTACAG	TTCCAACCTC	TTTAGCTTAC	AAAATGGCCA	GTCAAACGAT	TTTCTATCCT	1560
TTCATGCGAG	GGGATACTTA	TTTAAGAGTG	AACCCCTCAAT	TTGCAGTGCC	TAAAGGAACT	1620
GATGCTTCTAG	TCAGAGTGT	TGATGAATT	CATGGAAATG	CTTATTTAGA	AAATAACTAT	1680
AAAGTTGGTG	AAATCAAATT	ACCGATTCCG	AAATTAAAACC	AAGGAACAAAC	CAGAACGGCC	1740
GGAAAATAAAA	TTCCCTGTAAC	CTTCATGGCA	AATGCTTATT	TGGACAAATCA	ATCGACTTAT	1800
ATTGTGGAAG	TACCTATCTT	GGAAAAAAGAA	AATCAAACCTG	ATAAACCAAG	TATTCTACCA	1860
CAATTAAAAA	GGAAATAAAGC	ACAAGAAAAC	TCAAAACTTG	ATGAAAAGGT	AGAAGAACCA	1920
AAGACTAGTG	AGAAGGTAGA	AAAAGAAAAA	CTTTCTGAAA	CTGGGAATAG	TACTAGTAAT	1980
TCAACGTTAG	AGAAGTTCC	TACAGTGGAT	CCTGTACAAG	AAAAGTAGC	AAAATTGCT	2040
GAAAGTTATG	GGATGAAGCT	AGAAAATGTC	TTGTTTAAATA	TGGACGGAAC	AATTGAATTA	2100
TATTCTACCAT	CGGGGAGAAGT	CATTAAAAAG	AATATGGCAG	ATTTTACAGG	AGAAGCACCT	2160
CAAGGAAATG	GTGAAAATAA	ACCATCTGAA	AATGGAAAAG	TATCTACTGG	AACAGTTGAG	2220
AACCAACCAA	CAGAAAATAA	ACCAGCAGAT	TCTTTACCAAG	AGGCACCAAA	CGAAAAACCT	2280
GTAAAACCG	AAAACCTAAC	GGATAATGGA	ATGTTGAATC	CAGAAGGGAA	TGTGGGGAGT	2340
GACCCATGT	TAGATTCAAC	ATTAGAGGAA	GCTCCAGCAG	TAGATCTGT	ACAAGAAAAA	2400
TTAGAAAAAT	TTACAGCTAG	TTACGGATTA	GGCTTAGATA	GTGTTATATT	CAATATGGAT	2460
GGAACGATTG	AATTAAGATT	GCCAAGTGG	GAAGTGATAA	AAAAGAATT	ATTGATCTCA	2520
TAGCGTAA	(SEQ ID NO : 15)					2528

FIGURE 18

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CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDQVS QKEGIQAEQI	50
VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN	100
EVKGGYIIVK DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN	150
VAVARSQGRY TTNDGYVFNP ADIIIEDTGN A YIVPHGGHYH YIPKSDLSAS	200
ELAAAKAHLA GKNMQPSQLS YSSTPSPSLP INPGTSHEKH EEDGYGFDAN	250
RIIAEDESFGF VMSHGDHNHY FFKKDLTEEQ IKAQKHLLEE VKTSHNGLDS	300
LSSHEQDYP NAKEMKDLDK KIEEKIAGIM KQYGVKRESI VVNKEKNAAI	350
YPHGDHHAD PIDEHKPVGI GHSHSNYELF KPEEGVAKKE GNKVYTGEEL	400
TNVNLLKNS TFNNQNFTLA NGQKRVSF SF PPELEKKLG I NMLVKLITPD	450
GKVLEKVGK VFGEVGVNIA NFELDQPYLP GQTFKYTIA KDYPVSYDG	500
TFTVPTSLAY KMASQTIFYP FHAGDTYL RV NPQFAVPKG DALVRVDEF	550
HGNAYLENNY KVGEIKLPIP KLNQGTTRTA GNKIPVTFMA NAYLDNQSTY	600
IVEVPILEKE NQTDKPSILP QFKRNKAQEN SKLDEKVEEP KTSEKVEKEK	650
LSETGNSTSN STLEEVPTVD PVQEKAFA ESYGMKLENV LFNMDGTIEL	700
YLPGEVIIKK NMADFTGEAP QGNGENKPS NGKVSTGTVE NQPTENKPAD	750
SLPEAPNEKP VKPENSTDNG MLNPEGNVGS DPMLDSALEE APAVDPVQEK	800
LEKFTASYGL GLDSVIFNMD CTIELRLPSG EVIKKNLLIS	840
(SEQ ID NO : 16)	

FIGURE 19

CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDQVS QKEGIQAEQI	50
VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN	100
EVKGGYIIVK DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN	150
VAVARSQGRY TTNDGYVFNP ADIIIEDTGN A YIVPHGGHYH YIPKSDLSAS	200
ELAAAKAHLA GKNMQPSQLS YSSASDNNT QSVAKGSTSK PANKSENLQS	250
LLKELYDSPS AQRYSSESDGL VFDPAKIISR TPNGVAIPH DHYHFIPYSK	300
LSALEEKIAR MVPISGTGST VSTNAKPNEV VSSLGSLSSN PSSLTTSKEL	350
SSASDGYIFN PKDIVEETAT AYIVRHGDHF HYIPKSNQIG QPTLPNNSLA	400
TPSPSLPINP GTSHEKHEED GYGF DANRII AEDESFGVMS HGDHNHYFFK	450
KDLTEEQIKA AQKHLEEVKT SHNGLDSLSS HEQDYPGNK EMKDLDDKIE	500
EKIAGIMKQY GVKR EIVN KEKNAIIYPH GDHHADPID EH KPVGIGHS	550
HSNYELFKPE EGVAKKEGNK VYTGEELTNV VNLLKNSTFN NQNFTLANGQ	600
KRVSFSFPPE LEKKLG INML VKLITPDGKV LEK VSGKVFG EGVGNIANFE	650
LDQPYLPGQT FKYTIASKDY PEVSYDGTFT VPTSLAYKMA SQTIFYPFH	700
GDTYLRVNPQ FAVPKGTDAL VRV FDEFHGN AYLENNYKVG EIKLPIPKLN	750
QGTTRTAGNK IPVT FMANAY LDNQSTYIVE VPILEKENQT DKPSILPQFK	800
RNKAQENSKL DEKVEEPKTS EKVEKEKLSE TGNSTSNSTL EEPVTPDPVQ	850
EKVAKFAESY GMKLENVLFN MDGTIELYLP SGEV IKNMA DFTGEAPQGN	900
GENKPSENGK VSTGTVENQP TENKPADSLP EAPNEKPVKP ENSTDNGMLN	950
PEGNVGSDPM LDPALEEAAPA VDPVQEKLEK FTASYGLGLD SVIFNMDGTI	1000
ELRLPSGEVI KKNLSDFIA (SEQ ID NO : 55)	1019

FIGURE 20

CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDQVS QKEGIQAEQI 50  
 VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN 100  
 EVKGGYIIKV DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN 150  
 VAVARSQGRY TTNDGYVFNP ADIIIEDTGNA YIVPHGGHYH YIPKSDLSAS 200  
 ELAAAAKAHLA GKNMQPSQLS YSSTASDNNT QSVAKGSTSK PANKSENLQS 250  
 LLKELYDSPS AQRYSESDGL VFDPAKIISR TPNGVAIPHG DHYHFIPYSK 300  
 LSALEEKIAR MVPISGTGST VSTNAKPNEV VSSLGSLSSN PSSLTTSKEL 350  
 SSASDGYIFN PKDIVEETAT AYIVRHGDHF HYIPKSNQIG QPTLPNNSLA 400  
 TPSPSLPINP GTSHEKHEED GYGF DANRII AEDESGFVMS HGDHNHYFFK 450  
 KDLTEEQIKA AQKHLEEVKT SHNGLDSLSS HEQDYPGNA 489  
 (SEQ ID NO : 56)

FIGURE 21

MKFSKKYIAA GSAIVVSLSL CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDQVS 60  
 QKEGIQAEQI VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN 120  
 EVKGGYIIKV DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN VAVARSQGRY 180  
 TTNDGYVFNP ADIIIEDTGNA YIVPHGGHYH YIPKSDLSAS ELAAAAKAHLA GKNMQPSQLS 240  
 YSSTASDNNT QSVAKGSTSK PANKSENLQS LLKELYDSPS AQRYSESDGL VFDPAKIISR 300  
 TPNGVAIPHG DHYHFIPYSK LSALEEKIAR MVPISGTGST VSTNAKPNEV VSSLGSLSSN 360  
 PSSLTTSKEL SSASDGYIFN PKDIVEETAT AYIVRHGDHF HYIPKSNQIG QPTLPNNSLA 420  
 TPSPSLPINP GTSHEKHEED GYGF DANRII AEDESGFVMS HGDHNHYFFK KDLTEEQIKA 480  
 AQKHLEEVKT SHNGLDSLSS HEQDYPGNA 509  
 (SEQ ID NO : 57)

FIGURE 22

DLTEEQIKAQ KHLEEVKTS HNGLDSLSSH EQDYPGNAKE MKDLKKIEE 50  
 KIAGIMKQYG VKRESIVVNK EKNAAIYPHG DHHHADPIDE HKPVGIGHSH 100  
 SNYELFKPEE GVAKKEGNKV YTGEELTNVV NLLKNSTFNN QNFTLANGQK 150  
 RVSFSFPPPEL EKKLGINMLV KLITPDGKVL EKVSGKVFGE GVGNIANFEL 200  
 DQPYLPGQTF KYTIASKDYP EVSYDGTFTV PTSLAYKMAS QTIFYPFHAG 250  
 DTYLRVNPQF AVPKGTDALV RVFDEPHGNA YLENNYKVGE IKLPIPKLNQ 300  
 GTTRTAGNKI PVTFMANAYL DNQSTYIVEV PILEKENQTD KPSILPQFKR 350  
 NKAQENSKLD EKVEEPKTSE KVEKEKLSET GNSTSNTLE EVPTVDPVQE 400  
 KVAKFAESYG MKLENVLNFM DGTIELYLPS GEVIKKNMAD FTGEAPQGNG 450  
 ENKPSENGKV STGTVENQPT ENKPADSLPE APNEKPVKPE NSTDNGMLNP 500  
 EGNVGSDPML DPALEEAPAV DPVQEKEKF TASYGLGLDS VIFNMNDGTIE 550  
 LRLPSGEVIK KNLSDFPIAKL RYRSNHWVPD SRPEEPSPQP TPEPSPSPQP 600  
 APNPQPAPSN PIDEKLVKEA VRKVGDGYVF EENGVSRYIP AKNLSAETAA 650  
 GIDSKLAKQE SLSHKLGAKK TDLPSSDREF YNKAYDLLAR IHQDLDNKG 700  
 RQVDFEALDN LLERLKDVS DKVKLVDDIL AFLAPIRHPE RLGKPNAQIT 750  
 YTDDEIQVAK LAGKYTTEDG YIFDPRDITS DEGDAYVTPH MTHSHWIKKD 800  
 SLSEAERAAA QAYAKEKGLT PPSTDHQDSG NTEAKGAEAI YNRVKAACKV 850  
 PLDRMPYNLQ YTVEVKNGSL IIPHYDHYHN IKFEWFDEGL YEAPKGYTLE 900  
 DLLATVKYYV EHPNERPHSD NGFGNASDHV QRNKNGQADT NOTEKPSEEK 950  
 PQTEKPEEET PREEKPQSEK PESPKPTEEP EEESEPEESE PQVETEKVEE 1000  
 KLREAEDLQG KI QDPII KSN AKETLTGLKN NLLFGTQDNN TIMAEAEKLL 1050  
 ALLKESK (SEQ ID NO : 58)

FIGURE 23

CAYALNQHRS QENKDNNRVS YVDGSQSSQK SENLTPDQVS QKEGIQAEQI	50
VIKITDQGYV TSHGDHYHYY NGKVPYDALF SEELLMKDPN YQLKDADIVN	100
EVKGGYIIKV DGKYYVYLKD AAHADNVRTK DEINRQKQEH VKDNEKVNSN	150
VAVARSQGRY TTNDGYVFNP ADIIIEDTGNA YIVPHGGHYH YIPKSDLASAS	200
ELAAA (SEQ ID NO : 59)	205

FIGURE 24

CAYELGLHQA QTVKENNRVS YIDGKQATQK TENLTPDEVS KREGINAEQI	50
VIKITDQGYV TSHGDHYHYY NGKVPYDAII SEELLMKDPN YQLKSDIVN	100
EIKGGYVIKV NGKYYVYLKD AAHADNVRTK EEINRQKQEH SQHREGGTSA	150
NDGAVAFARS QGRYTTDDGY IFNASDIIED TGDAYIVPHG DHYHYIPKNE	200
LSASELAAAE AFLSGRENLS NLRTYRRQNS DNTPRTNWVP SVSNPGTTNT	250
NTSNNNSNTNS QASQSNIDS LLKQLYKLPL SQRHVESDGL IFDPAQITSR	300
TARGVAVPHG NYHFIPIYEQ MSELEKRIAR IIPLRYRSNH WVPDSRPEEP	350
SPQPTPEPSP SPQPAPNPQP APSNPIDEKL VKEAVRKVGD GYVFEENGVS	400
RYIPAKNLSA ETAAGIDSKL AKQESLSHKL GAKKTDLPPS DREFYNKAYD	450
LLARIHQDLL DNKGRQVDFE ALDNLLERLK DVSSDKVKLV DDILAFLAPI	500
RHPERLGKPN AQITYTDDEI QVAFLAGKYT TEDGYIFDPR DITSDEGDAY	550
VTPHMTHSHW IKKDSLSEAE RAAAQAYAKE KGLTPPSTDH QDSGNTEAKG	600
AAAIYNRVKA AKKVPLDRMP YNLQYTVEVK NGSLIIPHYS HYHNIKFEWF	650
DEGLYEAPKG YTLEDLLATV KYYVEHPNER PHSDNGFGNA SDHVQRNKNG	700
QADTNQTEKP SEEKPQTEKP EESTPREEKP QSEKPESPKP TEEPEEESPE	750
ESEEPQVETE KVEEKLREAE DLLGKIQDPI IKSNAKETLT GLKNLLFGT	800
QDNNTIMAEA EKLLALLKES K (SEQ ID NO : 60)	821

FIGURE 25

CAYELGLHQA QTVKENNRVS YIDGKQATQK TENLTPDEVS KREGINAEQI	50
VIKITDQGYV TSHGDHYHYY NGKVPYDAII SEELLMKDPN YQLKSDIVN	100
EIKGGYVIKV NGKYYVYLKD AAHADNVRTK EEINRQKQEH SQHREGGTSA	150
NDGAVAFARS QGRYTTDDGY IFNASDIIED TGDAYIVPHG DHYHYIPKNE	200
LSASELAAAE AFLSGRENLS NLRTYRRQNS DNTPRTNWVP SVSNPGTTNT	250
NTSNNNSNTNS QASQSNIDS LLKQLYKLPL SQRHVESDGL IFDPAQITSR	300
TARGVAVPHG NYHFIPIYEQ MSELEKRIAR IIPL	334
(SEQ ID NO : 61)	

FIGURE 26

RYRSNHWVPD SRPEEPSPQP TPEPSPSPQP APNPQPAPSN PIDEKLVKEA	50
VRKVGDGYVF EENGVSRYIP AKNLSAETAA GIDSKLAKQE SLSHKLGAKK	100
TDLPSSDREF YNKAYDLLAR IHQDLDNKG RQVDFEALDN LLERLKDVSS	150
DKVKLVDDIL AFLAPIRHPE RLGKPNAQIT YTDDEIQVAK LAGKYTTEDG	200
YIFDPRDITS DEGDAYVTPH MTHSHWIKKD SLSEAERAAA QAYAKEKGLT	250
PPSTDHQDSG NTEAKGAEAI YNRVKAAKKV PLDRMPYNLQ YTVEVKNGSL	300
IIPHEDHYHN IKFEWFDEGL YEAPKGYITLE DLLATVKYYV EHPNERPHSD	350
NGFGNASDHV QRNKNGQADT NQTEKPSEEK PTEKPEEET PREEKPQSEK	400
PESPKPTQEP EESPEESEE PQVETEKVEE KLREAEDLLG KIQDPIIKSN	450
AKETLTGLKN NLLFGTQDNN TIMAEAEKLL ALLKESK	487
(SEQ ID NO : 62)	

FIGURE 27

AEAFLSGRE	LSNLRTYRRQ	NSDNTPRTNW	VPSVSNP GTT	NTNTSNN SNT	50
NSQASQSNDI	DSLLKQLYKL	PLSQRHVESD	GLIFDPAQIT	SRTARGVAVP	100
HGNHYHPIY	EQMSELEKRI	ARIIPLRYRS	NHWVPSRPE	EPS PQPTPEP	150
SPSPQPAPNP	QPAPSNPIDE	KLVKEAVRKV	GDGYVFEENG	VSRYIPAKNL	200
SAETAAGIDS	KLAQESLSH	KLGAKKTDL	SSDREFYNKA	YDLLARIHQD	250
LLDNKG RQVD	FEALDNLLER	LKDVS SDKV	LVD DILAFLA	PIRHPERLGK	300
PNAQITYTDD	EIQVAKLAGK	YT TEDGYIFD	PRDITSDEGD	AYVTPHMTHS	350
HWIKKDSLSE	AERAAAQAYA	KEKGLTPPST	DHQDSGNTEA	KGAEAIYN RV	400
KA AKVPLDR	MPYNLQYTVE	VNGSLIIPH	YDHYHNKFE	WFDEGLYEAP	450
KGYTLEDLLA	TVKYYVEHPN	ERPHSDNGFG	NASDHVQRNK	NGQADTNQTE	500
KPSEEKPQTE	KPEEETPREE	KPQSEK PESP	KPTEEP EES	PESEEPQVE	550
TEKVEEKLR	AEDLLGKIQD	PIKSNAKET	LTGLKN NLLF	GTQDNNTIMA	600
EA EKLLALLK	ESK	(SEQ ID NO : 63)			613

FIGURE 28

DLTEEQIKAA	QKHLEEVKTS	HNGLDSLSSH	EQDYPGNAKE	MKDLDKKIEE	50
KIAGIMKQYG	VKRESIVVNK	EKNAAIIYPHG	DHHHADPIDE	HKPGVIGHSH	100
SNYELFKPEE	GVAKKEGNKV	YTGEELTNVV	NLLKNSTFNN	QNFTLANGQK	150
RVSFSFPPPEL	EKKLGINMLV	KLITPDGKVL	EKVSGKVFGE	GVGNIANFEL	200
DQPYLPGQTF	KYTIASKDYP	EVSYDGTFTV	PTSLAYKMAS	QTIFYPFHAG	250
DTYLRVNPQF	AVPKGTDALV	RVFDEFHGNA	YLENNYKVGE	IKLPIPKLNQ	300
GTTRTAGNKI	PVT FMANAYL	DNQSTYIVE	PILEKENQTD	KPSILPQFKR	350
NKAQENSKLD	EKVEEPKTSE	KVEKEKLSET	GNSTS NSTL	EVPTVDPVQE	400
KVAKFAESYG	MKLENVL FNM	DGTIELYLP	GEVIKKQNMAD	FTGEAPQGNG	450
ENKPSENGKV	STGTVENQPT	ENKPADSLPE	APNEKPVKP	NSTDNGMLNP	500
EGNVGSDPML	DPALEEAPAV	DPVQE KLEKF	TASYGLGLDS	VIFNMDGTIE	550
LRLPSGEVIK	KNLSDFIA	(SEQ ID NO : 64)			568

FIGURE 29

DLTEEQIKAA	QKHLEEVKTS	HNGLDSLSSH	EQDYPGNAKE	MKDLDKKIEE	50
KIAGIMKQYG	VKRESIVVNK	EKNAAIIYPHG	DHHHADPIDE	HKPGVIGHSH	100
SNYELFKPEE	GVAKKEGNKV	YTGEELTNVV	NLLKNSTFNN	QNFTLANGQK	150
RVSFSFPPPEL	EKKLGINMLV	KLITPDGKVL	EKVSGKVFGE	GVGNIANFEL	200
DQPYLPGQTF	KYTIASKDYP	EVSYDGTFTV	PTSLAYKMAS	QTIFYPFHAG	250
DTYLRVNPQF	AVPKGTDALV	RVFDEFHGNA	YLENNYKVGE	IKLPIPKLNQ	300
GTTRTAGNKI	PVT FMANAYL	DNQSTYIVE	(SEQ ID NO : 65)		329

FIGURE 30

EVPILEKENQ TDKPSILPQF KRINKAQENSK LDEKVEEPKT SEKVEKEKLS  
 ETGNSTSNT LEEVPTVDPV QEKVAKFAES YGMKLENVLF NMDGTIELYL  
 PSGEVVIKKNM ADFTGEAPQG NGENKPSENG KVSTGTVENQ PTENKPADSL  
 PEAPNEKPKV PENSTDNGML NPEGNVGSDP MLDPALEEAP AVDPVQEKL  
 KFTASYGLGL DSVIFNMDGT IELRLPSGEV IKKNLSDFIA  
 (SEQ ID NO : 66)

FIGURE 31

DIDSLLKQLY KLPLSQRHVE SDGLIFDPAQ ITSRTARGVA VPHGNHYHFI  
 PYEQMSELEK RIARIIPLRY RSNHWVPDSR PEEPSPQPTP EPSPSPQPAP  
 NPQPAPSNPI DEKLVKEAVR KVGDGYVFE NGVSRYIPAK NLSAETAAGI  
 DSKLAKQESL SHKLGAKKTD LPSSDREFYN KAYDLLARIH QDLDNKGRO  
 VDFEALDNLL ERLKDVSQDK VKLVDDILAF LAPIRHPERL GKPNAQITYT  
 DDEIQVAKLA GKYTTEDGYI FDPRDITSDE GDAYVTPHMT HSHWIKKDSL  
 SEAERAAAQA YAKEKGLTPP STDHQDSGNT EAKGAEAIYN RVKAAKKVPL  
 DRMPYNLQYT VEVKNGSLII PHYDHYNK FEWFDEGLYE APKGYTLEDL  
 LATVKYYVEH PNERPHSDNG FGNASDHVQR NKNGQADTNQ TEKPSEEKPQ  
 TEKPEEETPR EEKPKSEKPE SPKPTEEPEE ESPEESEEPQ VETEKVEKL  
 REAEDLLGKI QDPIIKSNAK ETLTGLKNL LFGTQDNNTI MAAEKLLAL  
 LKESK (SEQ ID NO : 67)

FIGURE 32

DIDSLLKQLY KLPLSQRHVE SDGLIFDPAQ ITSRTARGVA VPHGNHYHFI  
 PYEQMSELEK RIARIIPLRY RSNHWVPDSR PEEPSPQPTP EPSPSPQPAP  
 NPQPAPSNPI DEKLVKEAVR KVGDGYVFE NGVSRYIPAK NLSAETAAGI  
 DSKLAKQESL SHKLGAKKTD LPSSDREFYN KAYDLLARIH QDLDNKGRO  
 VDFEALDNLL ERLKDVSQDK VKLVDDILAF LAPIRHPERL GKPNAQITYT  
 DDEIQVAKLA GKYTTEDGYI FDPRDITSDE GDAYVTPHMT HSHWIKKDSL  
 SEAERAAAQA YAKEKGLTPP STDHQDSGNT EAKGAEAIYN RVKAAKKVPL  
 DRMPYNLQYT VEVKNGSLII PHYDHYNK FEWFDEGLYE APKGYTLEDL  
 LATVKYYVEH PNERPHSDNG FGNASDHV (SEQ ID NO : 68)

FIGURE 33

GLYEAPKGYT LEDLLATVKY YVEHPNERPH SDNGFGNASD HVQRNKNQQA  
 DTNQTEKPSE EKPQTEKPEE ETPREEPKQS EKPESPKPTE EPEEEESPEES  
 EEPQVETEKV EEKLRREAEDL L (SEQ ID NO : 69)

FIGURE 34

ASDHVQRNKN QADTNQTEK PSEEKPOTEK PEEETPREEK PQSEKPKESPK  
 PTEEEPEESP ESEEPQVET EKVEEKLREA EDLLGKIQDP IIKSNAKETL  
 TGLKNLLFG TQDNNTIMAE AEKLLALLKE SK  
 (SEQ ID NO : 70)

FIGURE 35

DIDSLLKQLY	KLPLSQRHVE	SDGLIFDPAQ	ITSRTARGVA	VPHGNHYHFI	50
PYEQMSELEK	RIARIIPLRY	RSNHWPDSR	PEEPSPQPTP	EPSPSPQPAP	100
NPQPAPSNPI	DEKLVKEAVR	KVGDGYVFE	NGVSRYIPAK	NLSAETAAGI	150
DSKLAKQESL	SHKLGAKKTD	LPSSDREFYN	KAYDLLARIH	QDLDNKGRQ	200
VDFEALDNLL	ERLKDVSSDK	VKLVDD	(SEQ ID NO : 71)		226

FIGURE 36

DILAFLAPIR	HPERLGKPNA	QITYTDDEIQ	VAKLAGKYTT	EDGYIFDPRD	50
ITSDEGDAYV	TPHMTHSHWI	KKDSLSEAER	AAAQAYAKEK	GLTPPSTDHQ	100
DSGNTEAKGA	EAIYNRVKAA	KKVPLDRMPY	NLQYTVEVKN	GSLIIPHVDH	150
YHNIKFEWFD	EGLYEAPKGY	TLEDLLATVK	YYVEHPNERP	HSDNGFGNAS	200
DHV	(SEQ ID NO : 72)				203

FIGURE 37

CSYELGRHQA	GQVKKESNRV	SYIDGDQAGQ	KAENLTPDEV	SKREGINAEQ	50
IVIKITDQGY	VTSHGDHYHY	YNGKVPYDAI	ISEELLMKDP	NYQLKDSIV	100
NEIKGGYVIK	VDGKYYVYLK	DAAHADNIRT	KEEIKRQKQE	HSHNHNNSRAD	150
NAVAARAQG	RYTTDDGYIF	NASDIIEDTG	DAYIVPHGDH	YHYIPKNELS	200
ASELAAAEEAY	WNGKQGSRPS	SSSSYNANPV	QPRLSEMHNL	TVTPTYHQNQ	250
GENISSLLRE	LYAKPLSERH	VESDGLIFDP	AQITSRTARG	VAVPHGNHYH	300
PIPYEQMSEL	EKRIARIIPL	RYRSNHWPVD	SRPEQPSPOS	TPEPSPSLQP	350
APNPQPAPSN	PIDEKLVKEA	VRKVGDGYVF	EENGVSRYIP	AKDLSAETAA	400
GIDSKLAKQE	SLSHKLGAKK	TDLPSSDREF	YNKAYDLLAR	IHQDLDNKKG	450
RQDFEVLDN	LLERLKDVSS	DKVVLVDDIL	AFLAPIRHPE	RLGKPNAQIT	500
YTDEIQVAK	LAGKYTTEDG	YIFDPRDITS	DEGDAYVTPH	MTHSHWIKKD	550
SLSEAERAAA	QAYAKEKGLT	PPSTDHQDSG	NTEAKGAEAI	YNRVKAAKKV	600
PLDRMPYNLQ	YTVEVKNGSL	IIPHVDHYHN	IKFEWFDEGL	YEAPKGYSLE	650
DLLATVKYYV	EHPNERPHSD	NGFGNASDHV	RKNKADQDSK	PDEDKEHDEV	700
SEPTHPESDE	KENHAGLNPSA	ADNLYKPSTD	TEETEEEAEAD	TTDEAEIPQV	750
ENSVINAKIA	DAEALLEKVT	DPSIRQNAME	TLTGLKSSLL	LGTKDNNNTIS	800
AEVDSLLALL	KESQPAPIQ	(SEQ ID NO : 73)			819

FIGURE 38

ENISSLLREL	YAKPLSERHV	ESDGLIFDPA	QITSRTARGV	AVPHGNHYHF	50
IPYEQMSELE	KRIARIIPLR	YRSNHWPDS	RPEQPSQST	PEPSPSLQPA	100
PNPQPAPSNP	IIDEKLVKEAV	RKVGDGYVFE	ENGVSRYIPA	KDLSAETAG	150
IDSKLAKQES	LSHKLGAKKT	DLPSSDREFY	NKAYDLLARI	HQDLDNKGR	200
QVDFEVLDNL	LERLKDVSSD	KVVLVDDILA	FLAPIRHPER	LGKPNAQITY	250
TDDEIQVAKL	AGKYTTEDG	IFDPRDITS	EGDAYVTPH	MTHSHWIKDS	300
LSEAERAAAQ	AYAKEKGLTP	PSTDHQDSGN	TEAKGAEAIY	YNRVKAAKVP	350
LDRMPYNLQY	YTVEVKNGSLI	IIPHVDHYHN	IKFEWFDEGLY	EAPKGYSLED	400
LLATVKYYVE	HPNERPHSDN	GFGNASDHVR	RKNKADQDSK	PDEDKEHDEVS	450
EPTHPESDEK	ENHAGLNPSA	DNLYKPSTD	EETEEEAEADT	TDEAEIPQVE	500
NSVINAKIAD	AEALLEKVTD	PSIRQNAME	TLTGLKSSLL	GTKDNNTISA	550
EVDSLLALLK	ESQPAPIQ	(SEQ ID NO : 74)			568

FIGURE 39

VRKNKADQDS KPDEDKEHDE VSEPTHPESD EKENHAGLNP SADNLYKPST  
 DTEETEEEAE DTTDEAEIPQ VENSVINAKI ADAEALLEKV TDPSIRQNM  
 ETLTGLKSSL LLGTDNNNTI SAEVDSLLAL LKESQPAPIQ  
 (SEQ ID NO : 75)

50  
 100  
 140

FIGURE 40

GAACAGACAG AAGAGCAAAT TAAGGCTGCG CAAAAACATT TAGAGGAAGT	50
TAAAACAGT CATAATGGAT TAGATTCTT GTCATCTCAT GAACAGGATT	100
ATCCAGGTA TGCCAAAGAA ATGAAAGATT TAGATAAAAA AATCGAAGAA	150
AAAATTGCTG GCATTATGAA ACAATATGGT GTCAAACGTG AAAGTATTGT	200
CGTGAATAAA GAAAAAAATG CGATTATTTA TCCGCATGGA GATCACCATC	250
ATGCAGATCC GATTGATGAA CATAAACCGG TTGGAATTGG TCATTCTCAC	300
AGTAACATATG AACTGTTAA ACCCGAAGAA GGAGTTGCTA AAAAAGAAGG	350
GAATAAAAGTT TATACTGGAG AAGAATTAAC GAATGTTGTT AATTGTTAA	400
AAAATAGTAC GTTAAATAAT CAAAACTTA CTCTAGCCAA TGTCAAAAA	450
CGCGTTCTT TTAGTTTCC GCCTGAATTG GAGAAAAAT TAGGTATCAA	500
TATGCTAGTA AAATTAATAA CACCAGATGG AAAAGTATTG GAGAAAGTAT	550
CTGGTAAAGT ATTTGGAGAA GGAGTAGGGA ATATTGCAAA CTTGATCAA	600
GATCAACCTT ATTTACCAGG ACAAAACATT AAGTATACCA TCGCTTCAAA	650
AGATTATCCA GAAGTAAGTT ATGATGGTAC ATTTACAGTT CCAACCTCTT	700
TAGCTTACAA AATGGCCAGT CAAACGATT TCTATCCTT CCATGCAGGG	750
GATACTTATT TAAGAGTGAA CCCTCAATT GCAGTGCCTA AAGGAACGTGA	800
TGCTTAGTC AGAGTGTGTT ATGAATTTCAT TGGAAATGCT TATTAGAAA	850
ATAACTATAA AGTTGGTGAA ATCAAATTAC CGATTCCGAA ATTAAACCAA	900
GGAACAAACCA GAACGGCCGG AAATAAAATT CCTGTAACCT TCATGGCAAA	950
TGCTTATTG GACAATCAAT CGACTTATAT TGTGGAAAGTA CCTATCTTGG	1000
AAAAAGAAAA TCAAACGTAT AAACCAAGTA TTCTACCCACA ATTAAAAGG	1050
AATAAAGCAC AAGAAAACCTC AAAACTTGTAT GAAAAGGTAG AAGAACCAAA	1100
GACTAGTGAG AAGGTAGAAA AAGAAAAACT TTCTGAAACT GGGAAAGTGA	1150
CTAGTAATTC AACGTTAGAA GAAGTTCCCTA CAGTGGATCC TGTACAAGAA	1200
AAAGTAGCAA AATTGCTGA AAGTTATGGG ATGAAGCTAG AAAATGTCTT	1250
GTAAATATG GACGGAACAA TTGAATTATA TTTACCATCA GGAGAAGTCA	1300
TTAAAAAGAA TATGGCAGAT TTTACAGGAG AAGCACCTCA AGGAAATGGT	1350
GAAAATAAAC CATCTGAAAA TGAAAAGTA TCTACTGGAA CAGTTGAGAA	1400
CCAACCAACA GAAAATAAAC CAGCAGATTC TTTACCAAGAG GCACCAAACG	1450
AAAAACCTGT AAAACCAAGAA AACTCAACCG ATAATGGAAT GTGAATCCA	1500
GAAGGGAAATG TGGGGAGTGA CCCTATGTTA GATCCAGCAT TAGAGGAAGC	1550
TCCAGCAGTA GATCCTGTAC AAGAAAAATT AGAAAATTT ACAGCTAGTT	1600
ACGGATTAGG CTTAGATAGT GTTATATTCA ATATGGATGG AACGATTGAA	1650
TTAAGATTGC CAAGTGGAGA AGTATAAAA AAGAATTAT CTGATTTCAT	1700
AGCGAAGCTT CGTTATCGTT CAAACCAATT GGTACCAAGAT TCAAGACCAG	1750
AAGAACCAAG TCCACAAACG ACTCCAGAAC CTAGTCCAAG TCCGCAACCT	1800
GCACCAAATC CTCACCAACG CTCAGCAAT CCAATTGATG AGAAAATTGGT	1850
CAAAGAAGCT GTTCGAAAAG TAGGCATGG TTATGTCCTT GAGGAGAATG	1900
GAGTTTCTCG TTATATCCCA GCCAAGAACATC TTTCAGCAGA AACAGCAGCA	1950
GGCATTGATA GCAAACCTGGC CAAGCAGGAA AGTTTATCTC ATAAGCTAGG	2000
AGCTAAGAAA ACTGACCTCC CATCTAGTGA TCGAGAATT TACAATAAGG	2050
CTTATGAGTT ACTAGCAAGA ATTCAACCAAG ATTTACTTGA TAATAAAGGT	2100
CGACAAAGTTG ATTTGAGGC TTGGATAAC CTGTTGGAAC GACTCAAGGA	2150
TGTCTCAAGT GATAAAAGTCA AGTTAGTGGA TGATATTCTT GCCTCTTAG	2200
CTCCGATTG TCATCCAGAA CGTTAGGAA AACCAAATGC GCAAATTACC	2250
TACACTGATG ATGAGATTCA AGTAGCCAAG TTGGCAGGGCA AGTACACAAC	2300
AGAAGACGGT TATATCTTG ATCCTCGTGA TATAACCAGT GATGAGGGGG	2350
ATGCCTATGT AACTCCACAT ATGACCCATA GCCACTGGAT TAAAAAGAT	2400

AGTTTGCTCG	AAGCTGAGAG	AGCGGCAGCC	CAGGCTTATG	CTAAAGAGAA	2450
AGGTTTGACC	CCTCCTTCGA	CAGACCATCA	GGATTCAGGA	AATACTGAGG	2500
CAAAAGGAGC	AGAACGCTATC	TACAACCGCG	TGAAAGCAGC	TAAGAAGGTG	2550
CCACTTGATC	GTATGCCTTA	CAATCTTCAA	TATACTGTAG	AAGTCAAAAA	2600
CGGTAGTTA	ATCATACCTC	ATTATGACCA	TTACCATAAC	ATCAAATTG	2650
AGTGGTTTGA	CGAAGGCCTT	TATGAGGCAC	CTAAGGGGTA	TACTCTTGAG	2700
GATCTTTGG	CGACTGTCAA	GTACTATGTC	GAACATCCAA	ACGAACGTCC	2750
GCATTCAGAT	AATGGTTTG	GTAAACGCTAG	CGACCATGTT	CAAAGAAAACA	2800
AAAATGGTCA	AGCTGATACC	AATCAAACGG	AAAAACCAAG	CGAGGGAGAAA	2850
CCTCAGACAG	AAAACCTGA	GGAAGAAACC	CCTCGAGAAG	AGAAACACAA	2900
AAGCGAGAAA	CCAGAGTCTC	CAAAACCAAAC	AGAGGAACCA	GAAGAAGAAT	2950
CACCAGAGGA	ATCAGAAGAA	CCTCAGGTCG	AGACTGAAA	GTTGAAGAA	3000
AAACTGAGAG	AGGCTGAAGA	TTTACTTGGA	AAAATCCAGG	ATCCAATTAT	3050
CAAGTCCAAT	GCCAAAGAGA	CTCTCACAGG	ATTTAAAAT	ATTTACTAT	3100
TTGGCACCCA	GGACAACAAT	ACTATTATGG	CAGAAGCTGA	AAAACATTG	3150
GCTTTATTAA	AGGAGAGTAA	G	(SEQ ID NO : 76)		3171

FIGURE 41

EAYWNGKQGS	RPSSSSSYNA	NPVQPRLSEN	HNLTVTPTYH	QNQGENISSL	50
LRELYAKPLS	ERHVESDGLI	FDPAQITSRT	ARGVAVPHGN	HYHFIPYEQM	100
SELEKRIARI	IPLRYRSNHW	VPDSRPEQPS	PQSTPEPSPS	LQPAPNPQPA	150
PSNPIDEKLV	KEAVRKVGDG	YVFEENGVSR	YIPAKDLSAE	TAAGIDSKLA	200
KOESLSHKLG	AKKTDLPSSD	REFYNKAYDL	LARIHQDLD	NKGRQVDFEV	250
LDNLLERLKD	VSSDKVKLVD	DILAFLAPIR	HPERLGKPNA	QITYTDDEIQ	300
VAKLAGKYTT	EDGYIFDPRD	ITSDEGDAYV	TPHMTHSHWI	KKDSLSEAER	350
AAAQAYAKEK	GLTPPSTDHQ	DSGNTEAKGA	EAIYNRVKAA	KKVPLDRMPY	400
NLQYTVEVKM	GSLIIPHVDH	YHNIFKFEWFD	EGLYEAPKGY	SLEDLLATVK	450
YYVEHPNERP	HSDNGFGNAS	DHV	(SEQ ID NO : 77)		473

FIGURE 42

CAYALNQHRS	QENKDNNRVS	YVDGSQSSQK	SENLTPDQVS	QKEGIQAEQI	50
VIKITDQGYV	TSHGDHYHYY	NGKVPYDALF	SEELLMKDPN	YQLKDADIVN	100
EVKGGYIIKV	DGKYYVYLKD	AAHADNVRTK	DEINRQKQEH	VKDNEKVNSN	150
VAVARSQGRY	TTNDGYVFNP	ADIIIEDTGNA	YIVPHGGHYH	YIPKSDLSAS	200
ELAAAKAHLA	GKNMQPSQLS	YSSTASDNNT	QSVAKGSTSK	PANKSENLQS	250
LLKELYDSPS	AQRYSESDGL	VFDPAKIISR	TPNGVAIPHG	DHYHFIPYSK	300
LSALEEKIAR	MVPISGTGST	VSTNAKPNEV	VSSLGSLSSN	PSSLTTSKEL	350
SSASDGYIFN	PKDIVEETAT	AYIVRHGDHF	HYIPKSNQIG	QPTLPNNSLA	400
TPSPSLPINP	GTSHEKHEED	GYGFDANRII	AEDESGFVMS	HGDHNHYFFK	450
KDLTEEQIKA	AQKHLEEVKT	SHNGLDSLSS	HEQDYPGNAK	EMKDLDKKIE	500
EKIAGIMKQY	GKVRESIVVN	KEKNAAIYPH	GDHHHADPID	EHKPGVIGHS	550
HSNYELFKPE	EGVAKKEGNK	VYTGEELTNV	VNLLKNSTFN	NQNFTLANGQ	600
KRVSFSFPPE	LEKKLGINML	VKLITPDGKV	LEKVGKVF	EGVGNIANFE	650
LDQPYLPGQT	FKYTIASKDY	PEVSYDGTFT	VPTSLAYKMA	SQTIFYPFHA	700
GDTYLRVNPQ	FAVPKGTDAL	VRVFDEFHGN	AYLENNYKVG	EIKLPIPKLN	750
QGTTRTAGNK	IPVTFMANAY	LDNQSTYIVE	(SEQ ID NO : 78)		780

FIGURE 43

CAYELGLHQA	QTVKENNRVS	YIDGKQATQK	TENLTPDEVS	KREGINAEQI	50
VIKITDQGYV	TSHGDHYHYY	NGKVPYDAII	SEELLMKDPN	YQLKDSDIVN	100
EIKGGYVIKV	NGKYYVYLKD	AAHADNVRTK	EEINRQKQEH	SQHREGGTS	150
NDGAVAFARS	QGRYTTDDGY	IFNASDIIED	TGDAYIVPHG	DHYHYIPKNE	200
LSASELAAA	AFLSGRENL	NLRTYRRQNS	DNTPRTNWV	SVSNPGTTNT	250
NTSNNNSNTS	QASQSNDIDS	LLKQLYKLPL	SQRHVESDGL	IFDPAQITSR	300
TARGVAVPHG	NHYHFIPYEQ	MSELEKRIAR	IIPLRYRSNH	WVPDSRPEEP	350
SPQPTPEPSP	SPQPAPNPQP	APSNPIDEKL	VKEAVRKVGD	GYVFEENGVS	400
RYIPAKNLSA	ETAAGIDSKL	AKQESLSHKL	GAKKTDLPSS	DREFYNKAYD	450
LLARIHQDLL	DNKGRQVDFE	ALDNLLERLK	DVSSDKVKLV	DDILAFLAPI	500
RHPERLGKPN	AQITYTDDEI	QVAKLAGKYT	TEDGYIFDPR	DITSDEGDAY	550
VTPHMTHSHW	IKKDSLSEAE	RAAAQAYAKE	KGLTPPSTDH	QDSGNTEAKG	600
AEAIYNRVKA	AKKVPLDRMP	YNLQVTVEVK	NGSLIIPH	HYHNIKFEWF	650
DEGLYEAPKG	YTLEDLLATV	KYYVEHPNER	PHSDNGFGNA		690

FIGURE 44

GTGAAGAAAA	CATATGGTTA	TATCGGCTCA	GTTGCTGCCA	TTTTACTAGC	TACTCATATT	60
GGAAGTTACC	AACTTGGTAA	GCATCATATG	GGTCTAGCAA	CAAAGGACAA	TCAGATTGCC	120
TATATTGATG	ACAGCAAAGG	TAAGGCAAAA	GCCCCCTAAAA	CAAACAAAAC	GATGGATCAA	180
ATCAGTGCTG	AAGAAGGCAT	CTCTGCTGAA	CAGATCGTAG	TCAAAATTAC	TGACCAAGGC	240
TATGTGACCT	CACACGGTGA	CCATTATCAT	TTTTACAATG	GGAAAGITCC	TTATGATGCG	300
ATTATTAGTG	AAGAGTTGTT	GATGACGGAT	CCTAATTACC	GTTTAAACA	ATCAGACGTT	360
ATCAATGAAA	TCTTAGACGG	TTACGTTATT	AAAGTCATG	GCAACTATTA	TGTTTACCTC	420
AAGCCAGGTA	GTAAGCGAA	AAACATTCGA	ACCAAACAAAC	AAATTGCTGA	GCAAGTAGCC	480
AAAGGAACTA	AAGAAGCTAA	AGAAAAAAGGT	TTAGCTCAAG	TGGCCCATCT	CAGTAAAGAA	540
GAAGTTGCAG	CAGTCATGA	AGCAAAAGA	CAAGGACGCT	ATACTACAGA	CGATGGCTAT	600
ATTTTTAGTC	CGACAGATAT	CATTGATGAT	TTAGGAGATG	CTTATTTAGT	ACCTCATGGT	660
AATCACTATC	ATTATATTCC	TAAGGAGGAT	TTGTCTCAA	GTGAGCTAGC	TGCTGCACAA	720
GCCTACTGGA	GTCAAAACAA	AGGTCGAGGT	GCTAGACCGT	CTGATTACCG	CCCGACACCA	780
GCCCCCAGGTC	GTAGGAAAGC	CCCAATTCCCT	GATGTGACGC	CTAACCCCTGG	ACAAGGTCTAT	840
CAGCCAGATA	ACGGTGGCTA	TCATCCAGCG	CTCCTAGGC	CAAATGATGC	GTCACAAAAC	900
AAACACCAAA	GAGATGAGTT	TAAGGAAAAA	ACCTTTAAGG	AACTTTTAA	TCAACTACAC	960
CGTCTTGATT	TGAAATACCG	TCATGTGGAA	GAAGATGGGT	TGATTTTGA	ACCGACTCAA	1020
GTGATCAAAT	CAAACGCTTT	TGGGTATGTG	GTGCTCATG	GAGATCATT	TCATATTATC	1080
CCAAGAAGTC	AGTTATCACC	TCTTGAAATG	GAATTAGCAG	ATCGATACTT	AGCTGGCCAA	1140
ACTGAGGACA	ATGACTCAGG	TTCAAGGAC	TCAAAACCAT	CAGATAAAGA	AGTGACACAT	1200
ACCTTTCTTG	GTCATCGCAT	CAAAGCTTAC	GGAAAAGGCT	TAGATGGTAA	ACCATATGAT	1260
ACGAGTGTG	CTTATGTTTT	TAGTAAAGAA	TCCATTCTT	CAGTGGATAA	ATCAGGAGTT	1320
ACAGCTAAC	ACGGAGATCA	TTTCACTAT	ATAGGATTG	GAGAACTTGA	ACAATATGAG	1380
TTGGATGAGG	TCGCTAACTG	GGTGAAGCA	AAAGGTCAAG	CTGATGAGCT	TGCTGCTGCT	1440
TTGGATCAGG	AACAAGGCAA	AGAAAAACCA	CTCTTGACA	CTAAAAAAAGT	GAGTCGCAA	1500
GTAACAAAAG	ATGGTAAAGT	GGGCTATATG	ATGCCAAAAG	ATGGTAAGGA	CTATTTCTAT	1560
GCTCGTGATC	AACTTGATTT	GACTCAGATT	GCCTTTGCCG	AACAAGAACT	AATGCTTAAA	1620
GATAAGAACG	ATTACCGTTA	TGACATTGTT	GACACAGGT	TTGAGCCACG	ACTTGTGTA	1680
GATGTGTCAA	GTCTGCCGAT	GCATGCTGGT	AATGCTACTT	ACGATACTGG	AAGTTGTTT	1740
GTTATCCCAC	ATATTGATCA	TATCCATGTC	GTTCGGTATT	CATGGTTGAC	GCGCGATCAG	1800
ATTGCAACAG	TCAAGTATGT	GATGCAACAC	CCCGAAGTTC	GTCCGGATGT	ATGGTCTAAG	1860
CCAGGGCATG	AAGAGTCAGG	TTCCGTATT	CCAAATGTTA	CGCCTCTTGA	AAAACGTGCT	1920
GGTATGCCAA	ACTGGCAAAT	TATCCATTCT	GCTGAAGAAG	TTCAAAAGC	CCTAGCAGAA	1980
GGTCGTTTG	CAACACCAGA	CGGCTATATT	TTCGATCCAC	GAGATGTTT	GGCCAAAGAA	2040
ACTTTTGAT	GGAAAGATGG	CTCCCTTGTG	ATCCCAAGAG	CAGATGGCAG	TTCATTGAGA	2100
ACCATTAAATA	AATCTGATCT	ATCCCAAGCT	GAGTGGCAAC	AAGCTCAAGA	GTATTGGCA	2160
AAGAAAAATA	CTGGTGATGC	TACTGATACG	GATAAACCCA	AAGAAAAGCA	ACAGGCAGAT	2220
AAGAGCAATG	AAAACCAACA	GCCAAGTGAA	GCCAGTAAAG	AAGAAAAAGA	ATCAGATGAC	2280
TTTATAGACA	GTTCACCGAGA	CTATGGTCTA	GATAGAGCAA	CCCTAGAAGA	TCATATCAAT	2340
CAATTAGCAC	AAAAGCTAA	TATCGATCCT	AAGTATCTA	TTTCCAACC	AGAAGGTGTC	2400
CAATTTTATA	ATAAAAATGG	TGAATTGGTA	ACTTATGATA	TCAAGACACT	TCAACAAATA	2460
AACCCTTAA	(SEQ ID NO : 80)					2469

FIGURE 45

VKKTYGYIGS	VAAILLATHI	GSYQLGKHHM	GLATKDNQIA	YIDDSKGKAK	50
APKTNKTMQ	ISAEEGISAE	QIVVKITDQG	YVTSHGDHYH	FYNGKVPYDA	100
IISEELLMTD	PNYRFKQSDV	INEILDGYVI	KVNGNYVYVL	KPGSKRKNIR	150
TKQQIAEQVA	KGTKEAKEKG	LAQVAHLSKE	EVAAVNEAKR	QGRYTTDDGY	200
IFSPTDIIDD	LGDAYLVPHG	NHYHYIPKKD	LSPSELAAAQ	AYWSQKQGRG	250
ARPSDYRPTP	APGRRKAPIP	DVTNPQGQGH	QPDNGGYHPA	PPRPNDAQN	300
KHQREDFKGK	TFKELLDQLH	RLDLKYRHVE	EDGLIFEPTQ	VIKSNAFGYV	350
VPHGDHYHII	PRSQLSPLEM	ELADRYLAGQ	TEDNDSGSEH	SKPSDKEVTH	400
TFLGHRIKAY	GKGLDGKPYD	TSDAYVFSKE	SIHSVDKSGV	TAKHGDHFHY	450
IGFGELEQYE	LDEVANWVKA	KGQADELAAA	LDQEQQKEKP	LFDTKKVSRK	500
VTKDGKVGYM	MPKDGKDYFY	ARDQLDLTQI	AFAEQELMLK	DKKHYRYDIV	550
DTGIEPRLAV	DVSSLPMHAG	NATTDTGSSF	VIPHIDHIHV	VPYSWLTRDQ	600
IATVKYVMQH	PEVRPDVWSK	PGHEESGSVI	PNVTPLDKRA	GMPNWQIIHS	650
AEEVQKALAE	GRFATPDGYI	FDPRDVLAKE	TFVWKDGGSFS	IPRADGSSLR	700
TINKSDLSQA	EWQQAQELLA	KKNTGDATDT	DKPKEKQQAD	KSNNENQQPSE	750
ASKEEKESDD	FIDSLPDYGL	DRATLEDHIN	QLAQKANIDP	KYLIFQPEGV	800
QFYNKNGELV	TYDIKTLQQI	NPP	(SEQ ID NO : 81)		823

FIGURE 46

GTGAAGAAAA	CATATGGTTA	TATCGGCTCA	GTTGCTGCCA	TTTTACTAGC	TACTCATATT	60
GGAAGTTACC	AACTTGGTAA	GCATCATATG	GGTCTAGCAA	CAAAGGACAA	TCAGATTGCC	120
TATATTGATG	ATAGCAAAGG	TAAGGCAAAA	GCCCCCTAAAA	CAAACAAAAC	GATGGATCAA	180
ATCAGTGCTG	AAGAAGGCAT	CTCTGCTGAA	CAGATCGTAG	TCAAAATTAC	TGACCAAGGT	240
TATGTGACCT	CACACGGTGA	CCATTATCAT	TTTTACAATG	GGAAAGTTCC	TTATGATGCG	300
ATTATTAGTG	AAGAGTTGTT	GATGACGGAT	CCTAATTACC	ATTTTAAACA	ATCAGACGTT	360
ATCAATGAAA	TCTTAGACGG	TTACGTTATT	AAAGTCAATG	GCAACTATTA	TGTTTACCTC	420
AAGCCAGGTA	GTAAGCGCAA	AAACATTCGA	ACCAAACAAAC	AAATTGCTGA	GCAAGTAGCC	480
AAAGGAACTA	AAGAAGCTAA	AGAAAAAGGT	TTAGCTCAAG	TGGCCCATCT	CAGTAAAGAA	540
GAAGTTGCGG	CAGTCATGA	AGCAAAAAGA	CAAGGACGCT	ATACTACAGA	CGATGGCTAT	600
ATTTTAGTGC	CGACAGATAT	CATTGATGAT	TTAGGAGACG	CTTATTAGT	ACCTCATGGT	660
AATCACTATC	ATTATATTC	AAAAAAAGAT	TTGTCTCCAA	GTGAGCTAGC	TGCTGCACAA	720
GCTTAUTGGA	GTCAAAAACA	AGGTGAGGT	GCTAGACCGT	CTGATTACCG	CCCGACACCA	780
GCCCCCAGGTC	GTAGGAAAGC	TCCAATTCT	GATGTGACGC	CTAACCCCTGG	ACAAGGTCTAT	840
CAGCCAGATA	ACGGTGGCTA	TCATCCAGCG	CCTCCTAGGC	CAAATGATGC	GTCACAAAAC	900
AAACACCAAA	GAGATGAGTT	TAAGGAAAAA	ACCTTTAAGG	AACTTTTACA	TCAACTACAC	960
CGTCTTGATT	TGAAATACCG	TCATGTGGAA	GAAGATGGGT	TGATTTTTGA	ACCGACTCAA	1020
GTGATCAAAT	CAAACGCTTT	TGGGTATGTG	GTGCCTCATG	GAGATCATT	TCATATTATC	1080
CCAAGAAGTC	AGTTATCACC	TCTTGAAATG	GAATTAGCAG	ATCGATACTT	AGCCGGTCAA	1140
ACTGAGGACA	ATGATTCAAG	TTCAGATCAC	TCAAAACCAT	CAGATAAAGA	AGTGACACAT	1200
ACCTTTCTTG	GTCATCGCAT	CAAAGCTTAC	GGAAAAGGCT	TAGATGGTAA	ACCATATGAT	1260
ACGAGTGATG	CTTATGTTT	TAGTAAAGAA	TCCATTCACT	CAGTGGATAA	ATCAGGAGTT	1320
ACAGCTAAC	ACGGAGATCA	TTTCCACTAT	ATAGGATTG	GAGAAGCTTGA	ACAATATGAG	1380
TTGGATGAGG	TCGCTAACG	GGTGAAGCA	AAAGGTCAAG	CTGATGAGCT	TGCTGCTGCT	1440
TTGGATCAGG	AACAAGGCAA	AGAAAAACCA	CTCTTTGACA	CTAAAAAAAGT	GAGTCGCAA	1500
GTAACAAAAG	ATGGTAAAGT	GGGCTATATT	ATGCCAAAAG	ATGGCAAGGA	CTATTTCTAT	1560
GCTCGTGATC	AACTTGATTT	GACTCAGATT	GCCTTTGCCG	AAACAGAACT	AATGCTTAA	1620
GATAAGAAC	ATTACCGTTA	TGACATTGTT	GACACAGGTA	TTGAGGCCAG	ACTTGCTGTA	1680
GATGTGTC	AACTGCGGAT	GCATGCTGGT	AATGCTACTG	ACGATACTGG	AAGTTGTTT	1740
GTTATCCCTC	ATATTGATCA	TATCCATGTC	GTTCCTGTATT	CATGGTTGAC	GCGCGATCAG	1800
ATTGCAACAA	TCAAGTATGT	GATGCAACAC	CCCGAAGTTC	GTCCAGATGT	ATGGTCTAAG	1860
CCAGGGCATG	AAGAGTCAGG	TTCGGTCTT	CCAAATGTTA	CGCCTCTTGA	TAACAGTGT	1920
GGTATGCCAA	ATTGGCAAAT	CATCCATTCT	GCTGAAGAAG	TTCAAAAGC	CCTAGCAGAA	1980
GGTCGTTTTG	CAACACCAGA	CGGCTATATT	TTCGATCCAC	GAGATGTTT	GGCCAAAGAA	2040
ACTTTTGAT	GGAAAGATGG	CTCCTTGTG	ATCCCAAGAG	CAGATGGCAG	TTCAATTGAGA	2100
ACCATTAATA	AATCTGATCT	ATCCCAAGCT	GAGTGGCAAC	AAGCTCAAGA	TTTATTGGCA	2160
AAGAAAAACG	CTGGTGATGC	TACTGATACG	GATAAAACCA	AAGAAAAGCA	ACAGGCAGAT	2220
AAGAGCAATG	AAAACCAACA	GCCAAGTGAA	GCCAGTAAAG	AAGAAGAAAA	AGAATCAGAT	2280
GACTTTATAG	ACAGTTTACC	AGACTATGGT	CTAGATAGAG	CAACCCCTAGA	AGATCATATC	2340
AATCAATTAG	CACAAAAAGC	TAATATCGAT	CCTAAGTATC	TCATTTCCA	ACCAGAAGGT	2400
GTCCAATTTC	ATAATAAAA	TGGTGAATT	GTAACCTATG	ATATCAAGAC	GCTTCACAA	2460
ATAAACCCCTT	AA	(SEQ ID NO : 82)				2472

FIGURE 47

VKKTYGYIGGS VAAILLATHI GSYQLGKHHM GLATKDQNQIA YIDDSSKGKAK	50
APKTNKTMDO ISAEEGISAE QIVVKITDQG YVTSHGDHYH FYNGKVPYDA	100
IISEELLMTD PNYHFKQSDV INEILDGYVI KVNGNYYVYL KPGSKRKNIR	150
TKQOIAEQVA KGTKEAKEKG LAQVAHLSKE EVAAVNEAKR QGRYTTDDGY	200
IFSPTDIIDD LGDAYLVPHG NHYHYIPKKD LSPSELAQQ AYWSQKQGRG	250
ARPSDYRPTP APGRRKAPIP DVTPNPGQGH QPDNGGYHPA PPRPNDASQN	300
KHQRDEFKGK TFKELLDQLH RLQLKVRHVE EDGLIFEPTQ VIKSNAFGYV	350
VPHGDHYHII PRSQLSPLEM ELADRYLAGQ TEDNDGSDH SKPSDKEVTH	400
TFLGHRIKAY GKGLDGKPYD TSDAYVFSKE SIHSVDKSGV TAKHGDHFHY	450
IGFGELEQYE LDEVANWVKA KGQADELAAA LDQEQQGKEKP LFDTKKVSRK	500
VTKDGKVGYI MPKDGKDYFY ARDQLDLTQI AFAEQEMLK DKNHYRYDIV	550
DTGIEPRLAV DVSSLPMHAG NATYDTGSSF VIPHIDHIHV VPYSWLTRDQ	600
IATIKYVMQH PEVRPDVWSK PGHEESGSVI PNVTPLDKRA GMPNWQIIHS	650
AEEVQKALAE GRFATPDGYI FDRPDVLAKE TFVWKDGFSFS IPRADGSSLR	700
TINKSDLSQA EWQQAQELLA KKNAGDATDT DPKKEKQQAD KSNENQQPSE	750
ASKEEEKESD DFIDSLPDYG LDRATLEDHI NQLAQKANID PKYLIFQPEG	800
VQFYNKNGEL VTYDIKTLQQ INPP (SEQ ID NO : 83)	824

FIGURE 48

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